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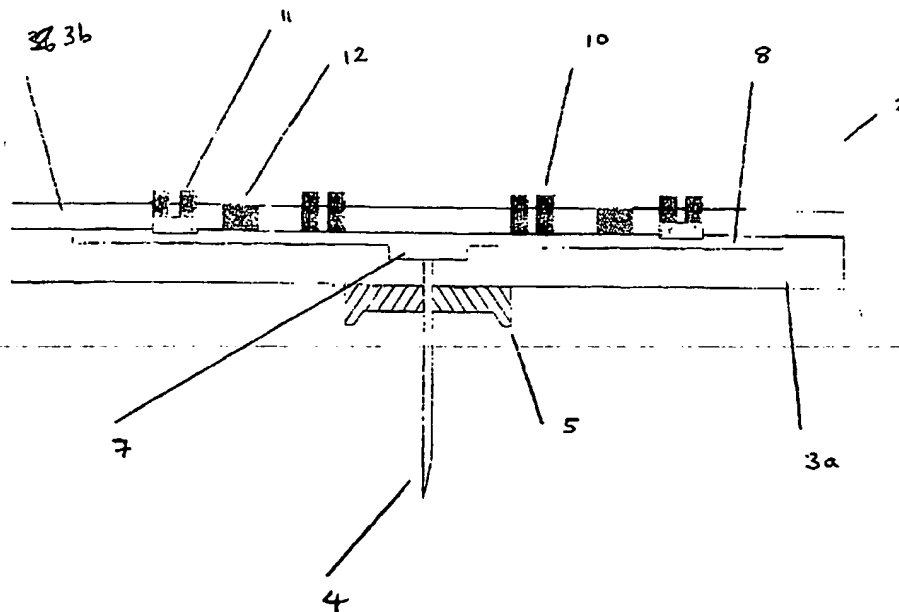
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[Continued on next page]

(54) Title: ANALYTE MEASUREMENT



(57) Abstract: A glucose sensor in the form of a skin patch (2) has a microneedle (4) which painlessly penetrates the skin to draw out interstitial fluid. The interstitial fluid passes to a common entrance port (7). A series of microchannels (8) is provided on the skin patch. The fluid drawn onto the patch is selectively switched between a number of microchannels (8) by means of electro-osmotic pumps (10) and hydrophobic gates (12). Each microchannel (8) has an electrochemical detector (11) for sensing glucose concentration. Also disclosed is a monolithic device with an integrated lance (83).

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## ANALYTE MEASUREMENT

## 5 I. Background of the Invention

This invention relates to apparatus for and methods of measuring certain properties of a fluid particularly although not exclusively, bodily fluids and the  
10 concentrations of certain analytes therein. At least some aspects of the invention relate particularly to the measurement of glucose levels in blood and other body fluids such as cutaneous (interstitial) and sub-cutaneous fluid.

15 Over recent years there have been two obvious trends in diagnostic device development: simplification of the test procedure and reduction in the volume of sample required to perform the test. Test simplification allows the assay to be performed by relatively untrained  
20 personnel in a "non-laboratory" setting. Thus, for example, cardiac marker tests configured in a lateral flow immunoassay format with labelled antibody allow for early assessment of potential myocardial infarct.

The driving force for the development of tests  
25 requiring smaller sample volumes is the reduction of discomfort of the patient. This is particularly important in home tests where, for example, if a glucose test is less painful the user will test more frequently. It is now well documented that diabetics who monitor  
30 their blood sugar frequently achieve better glycaemic control and so avoid long-term complications of the disease. With this thought in mind, a number of companies have developed test devices requiring progressively smaller sample volumes, thereby minimising  
35 pain.

Some proposals have been made to attempt to reduce

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the amount of pain involved by with so-called minimally invasive devices (e.g. Integ US patent numbers 5,746,217; 5,820,570; and 5,582,184). Unfortunately, such devices appear to be unable to provide sufficient fluid sample to provide a reliable result and thus, as yet, none has been commercially realised.

In the context of glucose measurement, so-called continuous monitors are also known and these have certain advantages over the 'snapshot' devices outlined above since they provide a clearer insight into trending, the effect of food or medication and overall glycaemic control. Such known devices however suffer from a number of drawbacks, mainly associated with the need to recalibrate the device regularly. This entails performing regular manual tests using a conventional test strip. This can negate many of the advantages of using a continuous device since the user is still relied upon to take action in order for the device to operate properly.

Furthermore the accuracy of the devices tends to drift unpredictably between calibrations. The regular finger pricks are also painful. Examples of continuous monitoring sensors include International patent application PCT/DE99/03126 (International Publication WO 00/22977 published 27 April 2000) and International patent application PCT/US99/16378 (International Publication WO 00/04832 published 3 February 2000).

## II. Summary and Objects of the Invention

~~It is an object of the invention to provide an~~  
improved arrangement and when viewed from a first aspect the invention provides a device for measuring the concentration of an analyte in a fluid, comprising a support member, analyte sensing means provided thereon for measuring said concentration and a microchannel in fluid communication with said analyte sensing means for conveying said fluid to said sensing means.

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Thus it will be seen that in accordance with the invention a microchannel is used to convey the sample fluid to the sensing means.

As used herein the term "microchannel" refers to a channel, of any suitable cross-section, whose lateral dimension is between approximately 10-500  $\mu\text{m}$ .

Such microchannels are beneficial for a number of reasons in the context of an analyte sensing device. In the context of the measurement of a bodily fluid a small sample volume is beneficial since it means that it is easier to provide a sufficient volume for a valid test. Most importantly the volume of fluid required to carry out an assay is correspondingly small, in addition the microchannel allows to handle flow rates between 10 and 500 nL/min with a high resolution of the measurement over time, due to the low channel volume. The analyte sensing means can be arranged in any suitable configuration depending on the type of sensing means used (electrochemical, optochemical etc.). Preferably the analyte sensing means comprises one or more reagents to react with said analyte and thereby give a measurable output. Such a reagent may be located anywhere, but preferably the reagent or at least one of the reagents is provided on a wall of the microchannel. This arrangement is beneficial since it means that no additional volume is required in order for the sample liquid to be brought fully into contact with the reagent - i.e. the contact area is optimised. When viewed from a second aspect the invention provides a device for measuring the concentration of an analyte in a fluid, comprising a support member and a microchannel provided on said support member for conveying said fluid across the support member, wherein one or more reagents for reacting with said analyte is coated on a wall of the microchannel.

Preferably such a device further comprises means for

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sensing the analyte - for example an electrochemical or a photometric detector.

The devices set out hereinabove may be used for measuring any suitable analyte in any suitable fluid.

5 However the invention, in its various aspects, finds particularly beneficial application in the measurement of analytes in human or animal bodily fluids. Of course, a suitable sample of bodily fluid may be applied to the device. Preferably however the device is integrated with

10 means for extracting the fluid. Most preferably this comprises means for penetrating the skin - in the case where the sample fluid is blood or interstitial fluid. Most preferably it comprises a skin penetration member such as hollow needle, solid lance or the like.

15 In accordance with the embodiments set out above the device can be used both to collect the sample fluid and to measure the concentration of the target analyte - or at least give an output from which the concentration can be measured. When viewed from a further aspect the

20 present invention provides a device for measuring the concentration of an analyte in a fluid, comprising a support member, means provided on the support member for sensing said analyte, wherein said support member further comprises fluid extraction means arranged to

25 direct fluid to said analyte sensing means via a conduit on the support member.

Thus it will be seen that in accordance with this aspect of the invention a fluid extraction means, preferably a skin penetration means, most preferably a

30 needle, lance or the like, is provided on a support member of a device for measuring analyte concentration.

This allows, in at least preferred embodiments, the combination of skin penetration means, e.g. needle, and support member, e.g. test strip, to be made disposable

35 and thus as hygienic as possible. The preferred embodiments require a user merely to puncture their skin

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with a needle or the like and then blood or interstitial fluid may be conducted automatically onto or into the test member. Devices for extracting interstitial fluid are shown in the afore-mentioned Integ patents and  
5 International patent application PCT/US01/09673 (International Publication WO 01/72220 published 4 October 2001).

In such embodiments the combined needle/test member would normally be used in conjunction with a separate,  
10 non-disposable, measuring device for measuring the analyte concentration e.g. by means of the output of an analyte-reagent reaction, where appropriate.

The dimensions of the needle, lance or the like may be suited to the particular application. In fact, any  
15 mechanism for extracting interstitial fluid may be used with the present invention. While the present invention will be described with reference to a shallow penetration needle, the invention is application to any technique for obtaining a body fluid.

20 For example a standard hypodermic needle several centimetres in length may be employed. Whilst the device may be used to sample interstitial fluid or blood from the cutaneous and/or the subcutaneous layers, in preferred embodiments however, the penetration member is  
25 of such a length that it penetrates substantially the cutaneous layer of skin. When the term "substantially the cutaneous layer" is referred to, it means that whilst it is the intention to sample fluid from the cutaneous layer, it cannot be ruled out that some sampling from  
30 the sub-cutaneous may occur. The cutaneous layer has a high density of blood capillaries which helps to ensure that the levels of analytes in interstitial fluid reliably reflect those in the blood.

While not limited to analysing interstitial fluid  
35 (ISF), the present invention is preferably used in obtaining and analysing ISF. A further benefit offered

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by interstitial fluid is that it is not as complex as whole blood and therefore does not, for example, suffer from Haematocrit fluctuations. ISF also has a low viscosity and is more suited to passage through  
5 microchannels than whole blood. The low sample volume requirement of a microchannel also enables the device reliably to use interstitial fluid.

Perhaps more importantly such shallow skin penetration (cutaneous or subcutaneous) substantially  
10 reduces the pain experienced since there is a much lower density of nerve endings than at the depths penetrated by standard needles when extracting blood. Indeed, in at least preferred embodiments, little or no pain is normally felt. This makes prolonged and/or repeated  
15 penetration of the skin much more acceptable.

The actual length of the penetration means will depend on the angle at which it is intended to be inserted. Thus if shallow angle penetration is intended, the length may be of the order of several millimetres  
20 e.g. up to approximately 7 to 10 millimetres. Preferably however substantially perpendicular insertion is intended and the length is less than 2 mm, preferably less than 1.5 mm. In some preferred embodiments a length of approximately 1.4 mm is appropriate. However for certain  
25 applications a length as short as 0.5mm may be preferred.

Of course a given test member may have just one skin penetration member, or alternatively a plurality may be provided.

The penetration means may comprise a sufficiently  
30 short standard needle, possibly shortened if necessary. More preferably the penetration means comprises a microneedle. A microneedle is hereby defined as a needle with a length sufficient to penetrate the cutaneous layer of human skin without substantially penetrating the sub-  
35 cutaneous layer. Such a needle typically has length less than 2mm, preferably between 0.4 and 1.6mm.



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The outer diameter is preferably less than 0.5mm, most preferably between 0.1 and 0.3 mm.

The penetration means need not be integrated with the support member, but preferably is. The comments  
5 above regarding this arrangement apply, with the additional benefit noted here that preferred embodiments of this arrangement can provide a compact disposable device which gives rise to substantially no pain, in use and obviates the need for a user to come into contact  
10 with or even see the bodily fluid concerned.

In a particularly preferred embodiment the analyte is glucose.

Preferred embodiments of the invention use a reagent-based measurement. Many different reagent-based  
15 analyte measurements are available to those skilled in the art, allowing the principles of the invention to find wide application. For example an optochemical- i.e. either fluorescent or luminescent - or an electrochemical technique could be used.

20 One preferred embodiment comprises analyte sensing means comprising a mediated amperometric enzyme electrode. For example, a ferrocene-mediated electron transfer from a glucose-oxidase catalysed reaction is a suitable means for detecting the level of glucose in a  
25 body fluid. Other enzymes, such as lactate oxidase or lactate dehydrogenase and cholesterol oxidase dehydrogenase may be used to measure lactate and cholesterol levels respectively. It will be appreciated that use of the above electron-transfer mediator is a  
30 non-limiting example and that other electron transfer mediators well known in the art could be used, for example, hexa-cyanoferrate (ferricyanide), oxygen or components of the respiratory chain (i.e. cytochromes) .

35 Preferably the embodiment will comprise an analyte sensing means which operates in conjunction with a test meter to give the measurement. Alternatively the member

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may comprise the appropriate reagent or dye for performing a colorimetric test in association with light sensitive means in the test meter. It will be seen that such arrangements are consistent with the test device  
5 being disposable since the costly elements of the sensing mechanism, such as the light sensitive means, electronic circuitry etc., can be placed in a non-disposable test meter.

Just single analyte sensing means may be provided,  
10 but preferably more than one is provided.

Thus it will be seen that in accordance with this aspect of the invention, a plurality of conduits direct fluid to be measured to respective analyte sensing means.

This means that a plurality of measurements of fluid may  
15 be made by a single such device. The sensing means may be different so as to measure or test for different analytes in the fluid. Preferably however they are the same or at least are for measuring the same analyte. When viewed from a further aspect the invention provides a  
20 device for measuring the concentration of an analyte in a fluid, comprising a support member and a plurality of analyte sensing means provided thereon for measuring said concentration, wherein said device further comprises a plurality of conduits such that each of said sensing  
25 means has a conduit associated therewith, said conduits serving in use to direct said fluid to respective sensing means. The device, whether it comprises a single or a plurality of conduit/analyte sensors, may be used to  
30 ~~continuously measure the concentration of the analyte~~  
over a period of time. Periodic measurements may be taken i.e. the fluid is "sampled" at various periods over time, for example every 30 minutes. Fluid may continuously flow through the device or may be temporarily stopped by, for example, hydrophobic gates.  
35 As yet another alternative, the sensors may be of the single measurement type, after which the device is

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discarded or fluid is diverted to another conduit to perform another measurement. The total time taken to complete a measurement cycle or the time taken between measurements could vary and would depend upon the degree of monitoring or the analyte of interest. The plurality of conduits would allow for switching from sensor to another after a particular period of time, by diverting fluid flow from one conduit to another. Switching enables measurements to be performed without interruption to the fluid extraction or without having to recalibrate the sensors, which are known to drift after a period of time due to factors such as electrode fouling etc.

Thus analyte monitoring may be made on a truly continuous basis, without the user having to recalibrate or interact with the system. An advantage of performing a single measurement, either in the case of a device with a single or plurality of microchannels is that drifting of the signal over time is no longer an issue that has to be considered and consequently simpler reagent chemistries may be employed in the analyte sensors. For example in the case of an electrochemical detection system, results may be obtained in as little as five seconds or less. Furthermore, such devices would not require storage reservoirs for collection of the waste fluid, since the waste fluid would be stored in the microchannel conduits themselves.

In principle, there would be no upper limit to the number of sensor/conduits contained within a device, the upper limit being determined by factors such as the desired total length of time for analyte measurement or by the size of the device.

Whilst the above embodiments describe a flow-through sensor, i.e. the analyte is measured whilst flowing past the detection means, measurements may also be conducted whilst the fluid is stationary. This could be achieved by the use of flow controlling means such as

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a hydrophobic gate or gates being activated to temporarily stop the flow of fluid through the microchannel once it had passed the sensing means. Isolating or shutting off a volume of fluid from the rest  
5 of the fluid sample would enable an end-point determination of the total amount of analyte present in the particular portion of the sample to be made.

As described above, the sensing means may be electrochemical or non-electrochemical in nature - e.g.  
10 of the fluorescent or chemi-luminescent colorimetric sort. For example the sensing means may comprises an enzyme-coated electrode in the case of electrochemical measurement, or in the case of fluorescent colorimetric sensing the sensing means would comprise a suitable  
15 reagent dye. It will, of course, be understood that the term 'sensing means' does not necessarily refer to a complete assembly for giving a final reading, but rather to a means on the support member which yields an output which may be read to give a measure of the analyte  
20 concentration, e.g. by a separate test meter.

Such devices according to embodiments  
of the invention as set out above could be arranged to be used in a mode similar to conventional test strips i.e. where a single fluid sample is placed on the device  
25 and is measured.

In common with earlier aspects of the invention, devices of the sort set out above may be arranged to measure the concentration of any suitable analyte. In all such cases, the analyte's concentration may simply be  
30 an indirect indication of the property of the sample fluid which it is desired to monitor. For example a detection reagent (e.g. an enzyme substrate or an antigen) may be added to the sample so as to bind selectively and/or react with certain proteins in the  
35 fluid such as enzymes or antibodies. In this case the added detection reagent or the product of the enzyme-

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catalysed reaction, comprises the analyte. Devices in accordance with the invention can therefore be arranged to measure such analytes - i.e. it will be seen that such devices may be used to measure the activity of enzymes in  
5 a body fluid sample, or test whether antibodies to a particular antigen are present in the body fluid.

Preferably however, the device is arranged to measure an analyte concentration directly - i.e. it is the concentration itself which is being monitored. An  
10 example of such a measurement would be glucose in blood, the concentration of which is an important parameter for those suffering from diabetes.

In a particularly preferred embodiment the measuring device is suitable for measuring the concentration of an  
15 analyte, e.g. glucose, in blood or interstitial fluid. In such an embodiment the device is preferably suitable for attachment to the skin of the subject who is to be measured. The subject may be an animal but is preferably human.

20 By providing a device which can be attached to a subject, measurements of the concentration of the substance in the subject's blood or interstitial fluid can be carried out repeatedly over a period of time.

This means that inconvenience and discomfort to the  
25 user is significantly reduced and moreover that tests can be carried out at regular intervals and at a greater frequency than would otherwise be tolerable. The result of this in the preferred application of the invention to  
~~blood glucose monitoring is that an improved insight into~~  
30 glycaemic control and the effects of food, medication and general trends on the glucose level is given. Moreover, by attaching the device to the patient and therefore making the device portable, it allows the patient to lead as normal life as possible. It also allows for the  
35 possibility of continuous measurements during periods when it may not be convenient to self-monitor by

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conventional manual methods of testing, for example during prolonged periods of exercise or whilst sleeping. The device could therefore include an alarm or means to activate an alarm in order to alert or wake the patient  
5 or any third party in case of a particular analyte level, for example in the case of hypoglycaemia.

Accordingly it is preferred that the devices comprise a penetration member with a sufficiently length substantially to penetrate the cutaneous layer of skin,  
10 most preferably integrally formed with the support member. It is further preferred that such devices comprises at least one microchannel for conveying the interstitial fluid to one or more of the sensing means.

When viewed from a further aspect the present  
15 invention provides a device for measuring the concentration of a given analyte in a bodily fluid comprising means for attaching the device to the skin of a subject and a plurality of sensing means for making a plurality of measurements of said concentration over a  
20 period of time.

It will be seen that effectively this aspect of the invention provides a device which can perform a plurality of measurements *in situ*, that is without user intervention being required. This has clear benefits in  
25 removing some constraints on the number, frequency and regularity with which measurements can be performed. Each sensing element in the device is designed to be used for a certain predetermined time during which the signal  
will not significantly drift over time. In this sense  
30 " significantly " represents the amount of drift permitted such the measurement result would not be affected by an clinically unacceptable amount. After such a period fluid is then switched to a new sensor and the process repeated. Preferably the device is provided with  
35 flow control means for influencing the flow of fluid to the sensing means. Any suitable flow control method may

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be employed with corresponding means to affect such a control method. For example Piezo-electric pumping, electrokinetic or mechanical methods such as 'unblocking' the flow along a selected conduit. - e.g. by  
5 allowing a gas bubble to escape or by opening a valve.

In certain embodiments the flow control means comprises a hydrophobic gate situated within the conduit/microchannel. A hydrophobic gate as herein disclosed refers to a hydrophobic surface region within a  
10 hydrophilic channel such that the flow of fluid is interrupted. By changing the hydrophobic nature of the gate, i.e. by making the hydrophobic region more hydrophilic, fluid may then be allowed to flow along the channel. Hydrophobic gates may be used to control the  
15 flow of fluid within a single microchannel or may be used to switch or redirect flow from one microchannel to another.

Alternatively, the hydrophobic nature of the gate may be maintained as it is and a increased pumping force  
20 (e.g. provided by a mechanical or electro-osmotic pump) may be applied in order for the fluid to breach the hydrophobic gate.

Preferably the device is arranged to direct fluid sequentially to each of the sensing means. The timing of  
25 the direction of the fluid to the sensing means could be pre-configured for example by software within the meter, such that the fluid is switched after a predetermined period of time. Preferably however the device comprises, or is adapted to interface with, control means to control  
30 said direction of the fluid to the sensing means. Such control is preferably automatic.

Additionally or alternatively however the control means may be such as to allow a user to specify when a measurement is to be made. This is beneficial as it  
35 allows measurement on demand which is useful for example in the case of blood glucose monitoring, as it allows the

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user to determine the effect on blood glucose of eating a particular snack or to determine how much insulin it is necessary to inject prior to eating or the user may simply want to carry out a check for reassurance. This enables the device to make periodic measurements of fluid from the body of the subject using fresh fluid for each measurement, thereby facilitating the desired object of monitoring the concentration of the substance in question over a period of time.

Where, as is preferred, each microchannel or other conduit is associated with a respective flow control means, each may be individually addressable by a suitable control means. This gives significant flexibility in how such a device may be used.

Preferably the device comprises a common fluid collection region in fluid communication with the bodily fluid to be measured - e.g. via a needle - each sensing means preferably being in selective fluid communication with said common collection region. Thus when viewed from a further aspect the present invention provides a device for making a plurality of measurements of the concentration of an analyte in a fluid comprising a common sample collection site in fluid communication with the fluid to be measured and a plurality of sensing means for measuring said concentration.

Where not otherwise specified, any suitable conduit may be provided for conveying the fluid to be measured to the sensing means, but preferred arrangements comprise ~~microchannels as defined herein.~~ Furthermore in the context of a plurality of analyte sensing means, the provision of microchannels allows many test elements to be positioned in close proximity thereby enabling even devices comprising a large number of sensing means to be made relatively small. This is beneficial in applications where the size and weight of the device is an important consideration, such as when it is attached



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to the user's body. .

As well as the understood benefits *per se* of a low sample volume arising from the use of microchannels, the Applicants have realised that such a small sample volume makes it practicable to change the mode of testing. More particularly it has been realised that a very low sample volume means that rather than conduct the usual reaction rate measurement as in known electrochemical devices e.g. for detecting blood analytes such as glucose, where electron transfer is measured as a function of time to determine the rate of transfer, an end-point test can be carried out in which the total amount of analyte in the sample volume is measured, thereby consuming substantially all of the analyte.

It has been appreciated that this is advantageous over rate measurements since the latter tend to be prone to interference, temperature and Haematocrit fluctuations (in the case of using blood). Whereas such a measurement method would take a prohibitively long time with known measuring devices, by using microchannels in accordance with the invention to give a required sample volume of the order of a few nano-litres, such a measurement can typically be completed in the order of a few seconds, thereby making it a practical proposition.

When viewed from a yet further aspect the present invention provides a method of measuring the amount of an analyte in a sample of liquid comprising providing an electrochemical measuring device having a sensor electrode within a microchannel, introducing said sample liquid into said microchannel, and measuring an aggregate current passed by said electrode to give an indication of said amount of analyte in the sample.

This aspect of the invention also extends to an electrochemical device for measuring the amount of an analyte in a sample of liquid comprising a sensor electrode disposed within a microchannel and means for

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measuring an aggregate current passed by said electrode in use to give an indication of said amount of analyte in the sample.

The invention also extends to an equivalent  
5 arrangement using a non-electrochemical sensing means - e.g. a colorimetric one.

Any suitable transfer means for transferring the fluid from the user's body to the device may be employed.

Preferably methods such as suction, ultra-sound or  
10 iontophoresis may be used. More preferably however the transfer means comprises a needle. In particularly preferred embodiments the needle is a microneedle as defined hereinabove.

The needle is preferably shaped to aid skin  
15 penetration. For example the tip region of the needle is preferably substantially conical. Furthermore it is preferred that the tip region has a reduced cross-section - preferably less than 0.2mm in width, most preferably less than 0.05mm in width.

20 Moreover the needle is preferably arranged to minimise the risk of blockage upon insertion into skin. For example the aperture of the needle may be provided on a side surface of the needle, rather than at the tip as is conventional. Preferably the aperture of the needle  
25 is recessed, thereby avoiding contact with the skin upon penetration and thus potential blocking and/or damage

The needle preferably has a bore such that the sample fluid is drawn up by capillary action. Suitable  
~~needle bore sizes range preferably from 21-30 most~~  
30 preferably 25. Any suitable inert and biocompatible material for the needle may be employed.

Examples of such inert materials include but are not limited to stainless steel, gold, platinum and metal-coated plastics

35 Most preferably one or more microneedles is employed as defined hereinabove.

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The methods above may be supplemented by applying pressure to the user's skin around the site at which it is penetrated. Such pressure could be applied purely manually, but preferably the device comprises means -  
5 e.g. suitably configured resilient means - to apply the pressure. The device may have skin pressurising means as explained in more detail below and as shown in the figures. Examples of skin pressurising means are set out in copending application US09/877,514 filed June 8, 2001.

10 Preferably therefore the devices of the preferred embodiments comprise display means to display the concentration of the analyte being measured such as blood glucose. Such display means may be coupled directly to the measuring device, but preferably it is  
15 separate from the device and receives data therefrom by telemetry. This approach has the advantage that the measuring device can be very light and thus comfortable to wear. For example the measuring device may be worn under clothing but monitored on a display means kept,  
20 say, in a pocket or for example in the form of a watch worn on the arm of the user, without the user having to disturb their clothing in order to view it.

Additionally or alternatively the device comprises or is coupled to means for administering a substance to a  
25 user on the device on the basis of the measured concentration. Thus in the previous example of a blood glucose monitoring device, rather than just being a passive monitoring device, undoubtedly useful *per se*,  
~~such a device can be used in conjunction with an insulin~~  
30 pump to maintain the user's glucose level within a desired range.

The insulin pump could be a separate device or alternatively could be integrated within the analyte sensing device *per se*. Controlling means present for  
35 example within the meter could control the amount of insulin administered by the insulin pump in response to

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the level of glucose measured by the device.

By effectively providing a feedback loop, preferred embodiments of this inventive feature can allow a diabetic, to maintain control of his/her glucose levels with a minimum of intervention i.e. only to replace consumable items such as a sensor/insulin patch.

Furthermore, particularly where a continuous sensing means is employed, tighter control of, say glucose level, is achievable than where intermittent manual tests and insulin administration are used.

Thus when viewed from a yet further aspect the present invention provides an apparatus for administering a substance to a user comprising a measuring device for measuring the concentration of an analyte in a bodily fluid from said user, said measuring device comprising a plurality of analyte sensing means and at least one conduit, preferably a microchannel, for conveying said bodily fluid to at least one of the sensing means; the apparatus further comprising means to administer said substance to said user on the basis of said measurement of concentration.

In accordance with all appropriate aspects of the invention, the measuring device preferably comprises a means for attachment to the skin of a user, for example in the form of a self-adhesive patch. This can provide a secure but comfortable arrangement for use over prolonged periods of time, and can be relatively unobtrusive.

The means for administering the substance may be ~~entirely separate from the measuring device or integrated~~ therewith. Preferably such an integrated administering means comprises a reservoir on or in the measuring device for dispensing the substance. In a particular embodiment the substance, such as insulin, is contained within a reservoir on an adhesive patch. The actual means for getting the substance from such a reservoir could comprises anything suitable such as a pressurised supply

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in conjunction with a flow control means such as a valve.

Preferably however a pump is used. In a preferred embodiment a single pump may be used both to administer the substance, such as insulin, to a user's body and also  
5 to draw out blood or preferably interstitial fluid, to a sensing means for making an analyte concentration measurement e.g. of glucose.

Thus, for example, a suitable device may comprise an adhesive patch comprising a microneedle (as defined  
10 herein) coupled to or in fluid communication with an array of microchannels and a separate needle for injecting insulin. In a preferred embodiment a single pump, e.g. a silicon micro-pump, may be used to inject insulin from a reservoir on the patch and to draw  
15 interstitial fluid over a glucose sensing means to a waste reservoir.

Where an embodiment of the invention calls for control and/or data processing means, these will generally comprise electronic means such as an integrated  
20 circuit or the like. A power supply is then also required. Preferably such control/processing means are portable. It or they may be provided in an integral package with the device e.g. the adhesive patch. Alternatively the control/processing means and/or power  
25 source may be provided separately and communicate with measuring device via a wire or wireless telemetry link as mentioned above.

The power source may be a battery or may be a 'renewable' source such as a solar cell or a dynamo  
30 energised by movement of the user. Of course a combination of these could be employed.

Measuring devices in accordance with the present invention may be fabricated using any suitable technique.

In particular, where provided, the microchannels may be  
35 made using any suitable micro-fabrication technique such as but not limited to embossing, plasma etching or

- 20 -

injection moulding

In one preferred embodiment, electrodes are provided on opposing sides of a fluid channel by forming a first channel which is filled with a conductive material, and  
5 forming a second channel for conveying the test fluid, the second channel cutting across the first thereby thereby forming two conductive portions within respective opposite sides of the second channel.

The conductive portions formed in accordance with  
10 the invention could be utilised for an electrochemical sensor arrangement. Micromachining techniques for making the abovementioned intersecting channels are preferred since they can be used to fabricate microchannels which can be formed close to one another, permitting dense  
15 arrays thereof. Furthermore, where the device has flow control means operating via electro-osmotic force, the driving electrodes are preferably positioned in close proximity to one another. This allows high electric fields to be achieved without  
20 applying unnecessarily high voltages. Preferably such driving electrodes are provided substantially on one side of a channel. In one embodiment the driving electrodes extend around the wall of the channel.

In a preferred variant of the method above, one or  
25 more electrodes may be formed on a second substrate which is then laminated to the main support member of the device. Methods used to deposit the electrodes onto the second substrate may be chosen preferably from a printing method, more preferably a screen-printing method.  
30 Alternatively, chemical or physical vapour deposition techniques could be employed. Generally speaking, the electrodes according to all embodiments of the invention may be formed of any suitable inert material such as carbon, gold, platinum, etc. According to one  
35 embodiment, carbon electrodes, optionally coated with reagents are provided on the second substrate by screen-

- 21 -

printing, which is then laminated onto the support member thus closing the channel or channels and two gold electrodes are provided adjacent one another on a substrate laminated onto the support member for an  
5 electro-osmotic pump.

If a second substrate is provided, preferably it is arranged to close the channel provided on the support member. This allows a very straightforward fabrication method in which electrodes are formed within a closed  
10 channel.

Lamination of one substrate to another will normally be carried out such that both laminates are perfectly aligned and that no further trimming or cutting is necessary. However, the device could be fabricated for  
15 example by firstly a lamination step followed by a cutting step whereby the second substrate may be trimmed to the shape of the support member. Lamination may be carried out by various methods such as ultrasonic or thermal welding or bonding, or by the use of an adhesive.

20 According to one embodiment, the support member is formed with an integral needle at one end and the second substrate is then laminated onto the support forming a channel and leaving the penetration member exposed. According to another embodiment the integrated skin  
25 penetration member is provided is open on one side. The penetration member is arranged so that upon insertion into skin, the skin itself effectively forms a wall of the member so that it can act like a hollow needle. Most preferably this is achieved by forming the penetration  
30 member with walls tapering away from the open side - e.g. a V shape. Thus when viewed from a further aspect the invention provides an apparatus for obtaining and measuring fluid, comprising a skin penetration member having at least one longitudinal side open, the other  
35 sides being arranged so as effectively to cause the penetrated skin to act as the remaining longitudinal side

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of the member when the penetration member is inserted into the skin.

Thus when viewed from a yet further aspect the present invention provides a method of fabricating a device for measuring the concentration of an analyte in a fluid comprising providing a support member, forming an open channel on a surface of the support member and laminating a second layer onto said support member so as to close said channel. The invention also extends to a device fabricated using such a method.

Where an optical measurement technique is employed, as an alternative to an electrochemical measurement technique a light sensing means will, in general, be required. In some cases a light source may also be required, but is not always the case, for example in the case of chemiluminescent measurement. Any such light sensing means and/or the light

source may be provided integrally with the non-disposable measuring device e.g. a test-meter. According to one embodiment it or they are provided separately of the part of the device which is brought into contact with the sample fluid - e.g. a skin patch. This means that the device itself can be made disposable while the relatively more expensive light sensing means and associated electronics for example could be provided in a separate test meter.

In preferred embodiments the test device comprises means for optimising the light transfer from the sensing means to the optically sensitive means. In a simple embodiment such means comprises a lens, e.g. integrally moulded as part of the support member for the test device. Additionally or alternatively the device may be arranged such that the light sensitive means views the sensing means along the conduit, e.g. microchannel, along which the sample fluid passes. In other words the



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conduit, preferably a microchannel, acts as a light pipe.

By measuring light transmission or reflectance along the microchannel rather than across it, the path length and therefore the optical density may be increased for a minimal sample volume. , thereby making its measurement easier and more accurate. The material from which the conduit is formed is preferably chosen so to maximise light throughput at the frequencies of interest.

It will be appreciated by those skilled in the art that the arrangement described above is beneficial in its own right in enhancing the signal that may be measured from a minimal sample volume and thus when viewed from a yet further aspect the present invention provides an apparatus for measuring the light from an assay comprising an elongate conduit portion along which a sample fluid is drawn in use and a light sensitive means arranged to be sensitive to light coming substantially from the longitudinal axis of said conduit portion.

In one particularly preferred embodiment a disposable skin patch is provided with a moulded plastics lens over the analyte sensing means. A corresponding test meter is designed to be placed over the patch and comprises a light sensitive element which sits over the lens when the meter is placed over the patch.

25

### III. Brief Description of the Drawings

Certain preferred embodiments of the invention will now be described, by way of example only, with reference to the accompanying drawings in which:

30

Figure 1 shows a first embodiment of the invention, in the form of a skin patch, in cross section;

35 Figure 2a shows an alternative microchannel arrangement;

- 24 -

Figure 2b shows a multiple channel/sensor arrangement;

Figure 3 depicts a cross section view of the skin patch  
of Figures 1, and 2 attached to the skin of the user with  
5 a controller unit attached to the skin patch;

Figure 4 depicts a display unit;

Figure 5 depicts another embodiment of the invention,  
10 showing schematically a skin patch integrated with a  
needle, which also acts as an electrode, in cross  
section;

Figure 6a depicts a microchannel and optochemical sensor  
15 in plan view;

Figure 6b depicts a microchannel and electrochemical  
sensor in plan view;

20 Figures 6c to e are cross sections of the microchannel of  
Figure 6b with fluid progressively entering the channel;

Figure 8a shows a further embodiment of the invention, in  
the form of a single-use device with an integrated  
25 puncturing means;

Figure 8b is a cross section through a user's skin having  
the device of Figure 8a therein;

30 Figures 8c and 8d depict the construction of a device  
similar to that in Figure 8a;

Figure 8e is an alternative embodiment, shown in  
perspective view, of a puncturing means;

35

Figure 8f is a still further embodiment of an

- 25 -

alternative puncturing means shown in perspective view;

Figure 8g is a side-sectional view of the embodiment of Figure 8f taken along line X - X in Fig. 8f;

5

Figure 8h is a cross-sectional view of the lance of Fig. 8e in tissue;

10 Figure 8i shows an integrally formed base member and lance showing a vent and a capillary sensing channel, but with upper laminate removed for clarity;

Figure 12a is a perspective view of a microchannel,

15 Figure 12b is a close-up, partial, perspective view of a microchannel; whereby electrodes are provided on opposing sides of a fluid channel by forming a first channel which is filled with a conductive material, and forming a second channel for conveying the test fluid, the second  
20 channel cutting across the first thereby thereby forming two conductive portions within respective opposite sides of the second channel.

25 Figure 17a is top view, taken in perspective, of a first preferred embodiment of a chip for extracting and distributing a fluid sample and having a meander shaped waste reservoir;

30 Figure 17b is the view of the embodiment of Figure 17a modified to illustrate a column shaped waste reservoir;

Figure 18 is a bottom view, taken in perspective, of the chip of Figure 17a;

35

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Figure 19 is a plan view of a measurement channel of the chip of Figure 17a;

Figure 20a is a plan view of the meander shaped waste reservoir of the embodiment of Figure 17a;

Figure 20b is a plan view of the column shaped waste reservoir of the embodiment of Figure 17b; and

Figure 21 is a plan view of the switching mechanism shown in Figure 20a.

#### IV. Detailed Description of the Drawings

##### A. Skin Patch - General Description

Turning to Figure 1 there is shown a skin patch 2 suitable for measuring the level of blood glucose in a user. Patch 2 is made up of two layers 3a and 3b

The lowermost layer 3a is made of an suitable microfabricated plastic such as polyester, polycarbonate, polystyrene, polyimide, or other suitable microfabricatable polymers of suitable dimensions to allow it to be worn comfortably for a prolonged period of time, and optionally has an adhesive on its underside to allow the patch to be securely attached to the skin of a user. An optional pressure ring (5) is designed to apply pressure to the surface of the skin to enhance the flow of fluid from the body of the patient into the device.

The pressure means may also be integrally formed and of the same material as that of 3a. The drawing is not to scale and thus the various features may be of a different relative thickness dimension than illustrated. It is understood that figure 1 is shown for illustrative purposes only and as such the underside of 3a may be of

- 27 -

any shape that would facilitate fixing of the device to the skin, for example when the pressure means is incorporated. The invention is not limited as such to a pressure ring and other types of pressure extraction designs could be used. A penetration device (4) e.g. a needle, lancet or cannula is attached to the lowermost layer 3a, through which fluid passes into the collection and fluid collection and distribution port (7) formed in the upper surface of layer 3a. Layer 3a has on its upper surface microfabricated channels. Port (7) is also formed by the same microfabrication process as the microchannels.

Laminated to the lowermost layer is a substrate layer 3b, preferably formed of the same material as that of the lowermost layer. This serves to close the microchannel. The microchannel and penetration member may optionally be coated with a hydrophilic material and/or an anti-coagulant such as a heparin attached to the inner surface such that during use it will not diffuse away. Also shown is the microchannel system 8, corresponding electrochemical detectors 11 and optionally the electro-osmotic pump systems 10 and the hydrophobic gates 12. Each of the electrochemical detectors 12 and electro-osmotic pump systems 10 are electrically connected via conductive tracks (not shown) to respective terminals (not shown) at the edge of the patch. Not shown in Figure 1 are the fluid reservoirs. Fig 2a shows another embodiment whereby hydrophobic gates are present on either side of the sensor.

With respect to Figure 3, a controller unit 102 is brought into contact with these terminals so as to make electrical contact in use in order to transmit the measurement output signals between the patch and a display unit 103 (see Figures 3 and 4).

A control unit 102 for the patch 2 is shown in Figure 3. As may be seen, this sits on top of the patch

- 28 -

2 and is secured to upper side of the patch in a suitable manner(not shown). The control unit serves to control the fluid pumping mechanism, hydrophobic gates, fluid switching means and analyte sensors. The control unit is  
5 also provided with means such that radio frequency communication may be made, for example with the meter, for transmission and receipt of data. The control unit would also determine when all the microchannel/sensors had been used and transmit a signal accordingly. The  
10 control unit would also have means to detect any malfunctioning of the device and to also transmit this information.

The control unit 102 is preferably battery operated and is capable of transmitting signals to a wireless  
15 display unit (Figure 4) for example a radio frequency communication.

Figure 2b shows a multichannel arrangement with the proximal end of the needle 4 is in fluid communication with a fluid collection and distribution port (7) formed  
20 in the substrate layer 3b of the patch. Radiating out from the manifold 6 are eighteen of the microchannel systems 8. Not shown is the fluid switching means.

#### B. Micro-Channel System with Flow Control

25 One such microchannel system 8 is shown in greater detail in Figure 6b. The microchannel 608 is in direct communication with the port 607 into which the fluid flows from the needle 4. Downstream of the port 607 is an electrode system comprising a pair of inert electrodes  
30 610 providing electrosmotic flow. The electrodes are formed on a second substrate layer 3b (omitted from this figure for clarity). The second substrate layer is then laminated onto the first substrate layer 3b thereby closing the microchannel 126 and bringing the electrodes  
35 610 into contact with it. The electrodes 610 are positioned adjacent to each other, together forming an

- 29 -

electro-osmotic pump system. By applying a voltage difference across the electrodes, an electric field is generated which drives fluid along the microchannel 608.

Positioned along the microchannel 608 is one or more  
5 hydrophobic gates 612 which prevents the flow of fluid by capillary action along the microchannel. Downstream of the hydrophobic gate 128 is an electrochemical sensor 611 comprising at least two electrodes also formed on the upper substrate layer 3b. At least one further  
10 hydrophobic gate similar to the first, is optionally provided downstream of the sensor 12. This serves to stop the flow of fluid past the electrodes to allow the possibility for an end-point detection of the analyte. It will be understood by those skilled in the art that  
15 the hydrophobic gates may be constructed of any suitable material, including, but not limited to, PTFE, polycarbonate, polyisobutylene, PMMA, dodecyl acetate, silicon rubber, synthetic wax, octadecyl mercaptane, dodecyl mercaptane, and/or octa decyltrichloro silane.

20

#### C. Skin Patch Operation

Operation of the skin patch 2 will now be described.

The patch is attached to the skin of a user causing the needle 4 to penetrate substantially into the cutaneous  
25 layer of skin. Cutaneous fluid is drawn up the needle 4 and into the port 7 of the device. Referring now to Figure 6c it will be seen that the fluid 136 is drawn into the microchannel 126 as far as the hydrophobic gate 128 where its flow is arrested. When a measurement is  
30 required, the controller unit 102 issues an appropriate signal to the electro-osmotic pump 610, in the form of a voltage difference across the two electrodes. This causes the fluid 136 to flow through the hydrophobic gate 128 and past the analyte sensor 611 until it reaches the  
35 second (optional) gate. Once the interstitial fluid 136 comes into contact with the sensor 12, a measurement is

- 30 -

performed by the analyte sensors.

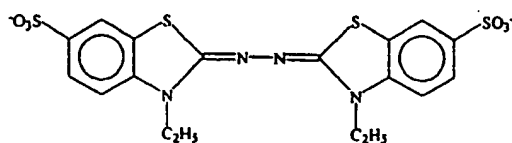
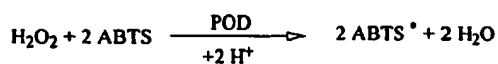
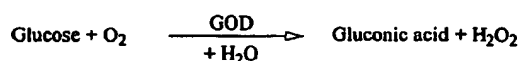
The controller unit 102 is provided with control means or means to initiate the flow of interstitial fluid sequentially to each of the microchannels (8) after a predetermined period of time. Once all the tests on the patch 2 have been performed, the controller 102 transmits a signal to the display means 103 to alert the user so that the patch 2 can be discarded and a fresh one applied.

10

#### D. Alternative Embodiment with Optochemical Sensing

An alternative form of microchannel 126 is shown in Fig. 6a. This is similar to that shown in Fig. 6b except that the electrochemical sensor 611 is replaced by an optochemical one 615. The optical sensor comprises a reaction chamber 616 which for example contains Glucose oxidase (GOD), Horseradish Peroxidase (POD) and a leuco-dye, (a colourless precursor of the dye molecule). (e.g. 2,2-Azino-di-[3-ethylbenzthiazoline-sulfonate]) The optical system is not limited to the above example and could comprise any suitable enzyme-dye combinations. The reaction which takes place is as below:

25 It will be seen from the above that the dye changes



Substrate / oxidized form of ABTS

colour in accordance with the amount of glucose in the sample. In order to measure this, the chamber 129 has an



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optically transparent upper surface. This arranged is to be aligned with the tip of an optical fibre in a modified control unit (not shown) so that the fibre is approximately 2 to 3 mm above the dye spot. The other  
5 end of the fibre is in optical contact with a light source and light sensitive diode sensor which is sensitive to a particular wavelength of light (for example 438nm in the case of ABTS) emitted by the transformed dye. The degree of absorption therefore  
10 gives a measure of the amount of transformed dye and thus the amount of glucose present.

As illustrated above, glucose present in the fluid acts as a substrate for GOD, which breaks the glucose down into glucono-1,5-lactone and hydrogen peroxide. POD  
15 subsequently catalyses the oxidation of the leuco-dye using the hydrogen peroxide, resulting in the production of a coloured dye which is subsequently detected. Those skilled in the art will understand that instead of 2,2-Azino-di-[3-ethylbenzthiazoline-sulfonate], any  
20 suitable leuco-dye may be used, e.g. Tetramethylbenzidine-Hydrochloride, or 3-Methyl-2-Benzothiazoline-Hydrazone in conjunction with 3-Dimethylamino-Benzoicacide.

The microchannels described as hereinbefore use  
25 electrochemical or colorimetric means to detect glucose.

The skilled person would understand that other detection means, such as infra red detection, filter photometry or Kromoscopy may be used.

### 30 F. Electrode Penetration Member

Figure 5 shows schematically a further preferred embodiment of the invention. similar construction to that of the previous embodiments and thus it has a hollow needle 106, for example but not limited to 1.4mm long and  
35 0.3mm wide which penetrates substantially the cutaneous layer of the skin 113 of the user..

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In this embodiment however the needle 106 also acts as an electrode which serves to extract fluid from the skin. At least one electrode 108 is provided such it makes contact with the skin. As shown in Fig 5, the  
5 needle is poised at a positive potential. By applying a voltage difference across the electrodes 106 and 108 an electric field is generated along the needle 106 and manifold 111 which stimulates the skin and increases the perfusion of interstitial fluid out of the skin.

10

#### G. Alternative Penetration Member Construction

Figures 8a to 8d show two further embodiments of the invention. In contrast with the foregoing embodiments, these are both single-use strips suitable for measuring  
15 blood glucose of a user. They are therefore used in a similar way to conventional test strips. However a principle difference is that each has an integrated lance 119 at one end.

Considering the device 115 shown in Fig 8a, it will  
20 be seen that it is made essentially of a layer 116 onto which a second layer is attached or laminated (not shown). The lowermost substrate layer 116 comprises a moulded or stamped microchannel 118, as well as the integrally formed lance 119 arranged in close proximity  
25 with the entrance 103 of the microchannel. During manufacture, the microchannel 118 maybe coated with suitable reagents, such as glucose oxidase in the case of glucose detection, which can be applied by any  
conventional means, such as printing for example screen-  
30 printing or ink-jet printing, or spray coating during manufacture. Alternatively, the microchannel may be free of enzyme, the reagents being provided on the electrode system (121) located on the underside of the uppermost layer (117).

35 The uppermost layer 117 may also be a conventional biosensor test-strip which could be attached to the lower

- 33 -

surface such that the electrodes are positioned on the underside of the formed channel. This particular electrode system 121 comprises three electrodes 221 of which at least one is covered with layers of enzyme and electron mediator such as glucose oxidase and ferricyanide to form a working electrode for detecting glucose, as is well known in the art, the other two being a counter and reference electrode. The electrode system may comprise two working electrodes which serve to compare currents at each and measure the channel filling speed of the blood or interstitial fluid. Alternatively the electrode system could comprise just two electrodes, a working and counter/reference. Not shown in Figures 1-21 is a venting device such that air may be displaced upon uptake of fluid into the microchannel. A vent or vents may be provided at any convenient location. Corresponding tracks 321 allow electrical connection to be made to the electrodes at the distal end of the strip when it has been inserted into a suitable test meter.

The upper layer 117 may be slightly longer than the lower one 116 to allow access to the tracks 321 for this purpose.

It will be noticed in particular that the lance 119 of the strip 115 in Fig 8a is essentially V shaped in cross section and tapers towards its tip. This means that when it is used to puncture a user's skin 123, as is shown in Figure 8b, the two sides of the V force back a portion of the skin 123, forcing the epidermis to form the remaining wall 123 of an enclosed channel 124. Thus an open channel is effectively transformed into a closed one when it is inserted into skin. This allows fluid to be drawn up the channel 124 so formed and into the microchannel 118, without having to mould a very fine hollow needle. The microchannel 118 may also be formed with a V shaped profile for convenience of fabrication, but this is not essential as may be seen from the

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slightly modified embodiment of Figs 8c and 8d in which the microchannel 118' has a rectangular profile.

In the use of the strip 115, the user pierces their skin with the lance 119 and interstitial fluid or blood is made to flow, by means of capillary action, through channel 124 formed by the lance (119) and the skin 122, into the microchannel 118 and thus over the electrochemical detector 121.

The user then withdraws the strip 115 and inserts the opposite end into a conventional style test meter which makes electrical connection to the three conductive tracks 321. Preferably however, the meter is an integral part of the lancing system such that the user does not need to insert the strip into the meter and is automatically provided with the measurement result.

Figures 8e through 8h show alternative embodiments of lances for penetrating into a body-fluid laden layer of skin as an alternative to the embodiment of the lance 119 of Figs. 8a and 8b. In Figure 8e, the lance 119a is an integrally formed pointed protrusion from the device 115 (not shown in Figs. 8e but identical to that in Fig. 8a) with a longitudinal capillary channel 121a cut completely through the thickness of the lance 119a. At a pointed distal tip 125a of the lance 119a, the lance 119a is provided with an enlarged area 123a of the channel 121a. The enlarged area 123a also is cut completely through the thickness of the lance 119a. At its proximal end 127a, the capillary channel connects with the microchannel 118 of the device 115 of Fig. 8a.

As shown best in Fig. 8h, the embodiment of Fig. 8e permits fluid to enter into the capillary channel 121a from opposite sides of the lance 119a and with the wall of the skin cooperating with the walls of the lance 119a to define an enclosed channel 121a. Fluid then can accumulate in the pooling area 123a and flow from the pooling area 123a into the capillary channel 121a as well

- 35 -

as flow directly from the skin into the capillary channel 121a for passage to the microchannel 118.

In the embodiment of Figure 8f, a lance 119b is of a design similar to that of Fig. 8e is shown but excluding  
5 the large pooling area 123a. Elimination of the pooling area 123a permits a narrower transverse dimension to the lance 119b.

In addition to fabricating the device from molded parts, the base member 116 and lances 119, 119a, 119b can be  
10 stamped from electrically conductive material. In such cases, the base member may be an electrode. An electrically conductive base member and lance can be stamped from metal as described or formed in any other acceptable manner (e.g., photochemically etching a metal  
15 stock material, machining or other fabricating technique). While the electrically conductive base member can be made of stainless steel it can be also be plated with a second metal such as for example gold, platinum or silver. The electrically conductive base  
20 material may also be used as a counter electrode.

Fig 8i shows an integrally formed base member and lance stamped from preferably one piece of sheet metal. The metal is preferably, but not limited to, stainless steel optionally coated with a noble metal such as gold  
25 or silver. Also shown on the base sheet is a microchannel 84 onto which a second layer such as a test-strip could be attached. It is intended that such a device as illustrated in Fig 8i would be incorporated  
into a lancet firing device such that the device could be  
30 fired into the skin. Furthermore the individual devices may be loaded and stored in a cassette comprising individually sealed pouches. Such a cassette could also be stored within the lancet firing device.

The stamped penetration member with a rectangular  
35 vent 81 which also serves as a capillary break ensuring that once fluid is taken up by the lance 83 into the

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sensing zone 82, the flow of fluid is halted. The vent may be of any suitable size or shape.

#### J. Detailed Preferred Embodiment

##### 5        1. Sampling Module

With reference now to Figures 17a - 21, the present invention will now be described with reference to a most preferred embodiment of the invention in an ex vivo continuous monitoring system.

10        The ex vivo continuous monitoring system consists of several major subsystems including sample extraction, sample fluid distribution, and electrochemical detection. As will be more fully described below, a body fluid sample can be extracted from the skin and guided into a  
15        channel for the electrochemical determination of glucose concentration.

After a predetermined period, the body fluid is directed into a different, previously unused channel. This process avoids false results otherwise arising from  
20        fouled electrodes or denaturant enzymes. Since the channels into which the fluid is re-directed are not filled with a sample solution or any other liquid, the electrochemical system in the channel will not show aging effects.

25        The disposable portion of the ex vivo continuous monitoring system is referred to herein as a chip. Figures 17a, 17b show embodiments of such a chip 400, 400a. In the embodiments of Figures 17a, 17b, each chip 400, 400a has for example, but not limited to, twelve  
30        measurement channels 402, 402a with switching mechanism 404 and 404a and waste reservoirs 406, 406a. Differences in the embodiments of chips 400, 400a will be described below with initial description being limited to a discussion of chip 400. It will be appreciated that chip  
35        400a is identical to chip 400 except as will be described. Similarly elements are numbered similarly

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with the addition of an "a" to distinguish embodiments.

The chip 400 has an edge connector 401 for electrically connecting electrodes (as will be described) to a controller (not shown). A controller can contain  
5 logic, memory and processors for controlling the electronics of the chip 400 and optionally displaying any outputs (as, for example, the function and operation of controller 102 in Fig. 3).

Preferably the measurement results are transmitted  
10 by radio frequency by the controller to a remote control allowing the user to view the result and optionally to communicate or interact with the controller of the monitoring system.

The sampling chip 400 extracts body fluid from the  
15 skin. In the preferred embodiment, the chip 400 extracts interstitial fluid (ISF) from substantially the cutaneous layer of a patient. With reference to Fig. 18, the chip 400 includes a suitable cannula or needle 410, a spring-loaded hub 412 to pressurize the skin and means such as  
20 an adhesive 414 to fix the chip 400 to the skin. Sampling modules with needles and spring-loaded hubs are shown in U.S. Pat. Nos. 6,203,504 and 5,746,217. It will be appreciated the foregoing description is a preferred embodiment. Any technique for obtaining a sample of body  
25 fluid may be used in combination with the teachings of the present invention.

Body fluid enters the needle 410 due to the pressure applied by the hub 412 and travels in to the micro-channels 402 as described above with reference to Figs.  
30 1a, 1b and 2. The needle 410 discharges the body fluid into a common distribution depression 416 shown best in Fig. 19). The distribution depression 416 is in fluid flow with the micro-channels 402. In the embodiment shown (having 12 channels of the dimensions described,  
35 the shape and the volume to the distribution depression

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416 is preferably a disk shaped void with 1 mm diameter and 100  $\mu\text{m}$  depth. These dimensions and shape may vary depending upon the size, number and flow rate of individual channels. Not shown in the drawings, a  
5 portion of the micro-channel 402 between the distribution depression 416 and a ground or reference electrode 420 may be restricted in size to reduce flow rate variability and therefore control the rate of flow.

Means may be provided within the device for the  
10 removal of undesirable gas-bubbles from the fluid sample. Such means could comprise a filter positioned between the needle and the common reservoir.

## 2. Continuous Electrochemical Detection of Glucose

15 The electrochemical detection system consists of an electrode system shown best in Fig. 19. The electrode system includes at least one working electrode 418 (by way of non-limiting example, gold or carbon) and a reference electrode 420 (e.g., silver/silver-chloride).  
20 The reference electrode 420 is circular to serve as a common electrode for each channel 402, thus reducing the number of contacts necessary. Alternatively, each channel can be provided with a unique reference electrode. Each channel 402 has a unique dedicated  
25 working electrode 418. In addition, a counter electrode (not shown) may be added. The electrodes 418, 420 are disposed to contact fluid in the channel 402. As previously described, the channels 402 are formed as open channels in a substrate layer. The electrodes 420,  
30 418 are screen-printed on an overlying laminate layer which also serves to seal the channels 402. The working electrode material is dependent on the enzyme ink being used. For example, one can distinguish between a redox-mediated system for the carbon working electrodes and an



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oxygen mediated system for the gold or platinum electrodes.

While a specific electro-chemical system will now be described, it will be appreciated that such description  
5 is illustrative and any suitable system could be used.

In the case of the redox-system, an organo-metal-complex (which is linked to polymers forming the ink body) acts as a conductor and electron acceptor for the enzyme. The electrons are transported by the redox-  
10 mediator directly or by an electron hopping mechanism from redox-center to redox-center to the working electrode. At the working electrode, the mediator is finally recycled to its original redox-state before it can react with the enzyme protein in a new cycle. The  
15 following are suitable for this detection method: glucose oxidase from aspergillus niger (GOD EC 1.1.3.4) and PQQ dependent glucose dehydrogenase (GDH EC 1.1.99.17).

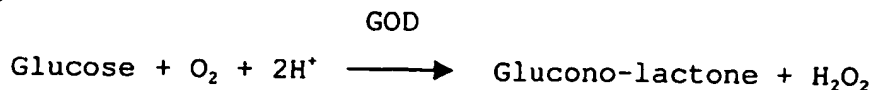
The enzyme ink is a cross-linked hydrophilic gel allowing the penetration of sample fluid but restricting  
20 the movement of the enzyme thus it is fixed to the electrode. This is an important property of the enzyme ink because the concentration of the enzyme has an effect upon the total response of the sensor. The same effect can be observed with the redox-mediator, due to the small  
25 molecular weight it cannot be entrapped in the same way as enzymes. Therefore, in the preferred embodiment, it is preferred to link the redox-mediator molecule to the large molecules such as the enzymes, proteins, or to the polymers of the ink material directly. In case of oxygen-  
30 mediated systems the situation is simpler. The enzyme ink needs only to contain enzyme as active component, the oxygen itself is dissolved in the sample fluid and is transferred to the sensor from the sample stream in the same instance as glucose.

35 In case of glucose determination the oxygen-mediated system can be created with GOD based on the

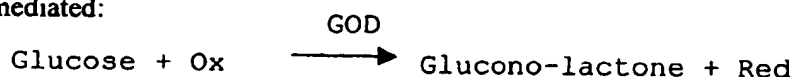
- 40 -

ability of the enzyme to build hydrogen peroxide from oxygen and glucose.

5 Oxygen-mediated:



10 Redox-mediated:

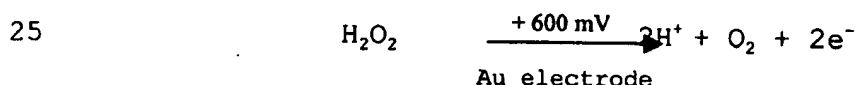


15 Ox = oxidized mediator species e.g.  $\text{K}_3[\text{Fe}(\text{CN})_6]$ , p-benzoquinon, Ferrocinium

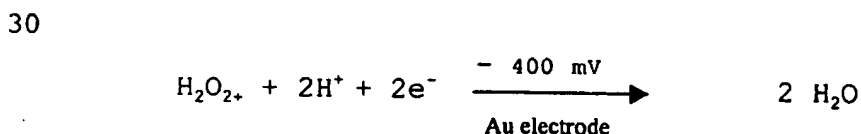
Red = reduced mediator species e.g.  $\text{K}_4[\text{Fe}(\text{CN})_6]$ , hydroquinone, Ferrocene

20 Hydrogen peroxide is a very reactive molecule, which can function as mediator to transport electrons for the enzyme to the electrode surface.

Oxidation of hydrogen peroxide on gold:



Reduction of hydrogen peroxide on gold:



35 Normally only the oxidation of hydrogen peroxide is used for analytical purposes due to the possibility of direct oxygen reduction on negative polarized noble metal electrode during the reduction of hydrogen peroxide. However, the reduction of hydrogen peroxide would open the possibility to use more cost effective electrode materials such as silver instead of gold, platinum, or palladium. In comparison to the noble metal electrodes we need for the oxidation or reduction

40

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of hydrogen peroxide carbon electrodes can be used for the oxidation of the redox-mediator.



In the preferred embodiment, the electrodes are deposited on a film such as polystyrene or polycarbonate of about 10 - 200  $\mu\text{m}$  and, more preferably, about 30 to 75  $\mu\text{m}$ ) using screen-printing technology. Subsequently to this print step the working electrode is coated with ink containing enzyme and in the case of carbon working electrodes with an ink containing enzyme and mediator.

This coating step could be accomplished with the same printing methods as used for the electrode material. Thus the production could be done in one machine with different print heads or stations on a continuous web. Such a process is described in "Continuous Process for Manufacture of Disposable Electrochemical Sensor" , US patent application Ser. No. 09/537599.

25            3. Fluidic Elements of Ex Vivo Continuous  
Monitoring

A schematic diagram of one measurement channel 402 is shown in Fig. 19. The channel 402 extends from the needle 410 and distribution depression 416 and through the main channel 402 (of dimensions for example of 200  $\mu\text{m}$  x 100  $\mu\text{m}$ ) containing the electrodes 418, 420.

As previously described, a flow restricted portion of the channel 402 namely a small cross section (e.g., 30  $\mu\text{m}$  x 30  $\mu\text{m}$ ) may be provided between the distribution 35 depression 416 and the common electrode 420. Such a flow restriction levels out different flow rates during the extraction of the body fluid. These flow rate differences

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are caused due to the changing physiological conditions during the extraction. In general, one could observe a higher flow rate at the beginning of a sampling period compared to the extraction rate at the end. A thin  
5 capillary can counter-act this behavior.

The electrodes 418, 420 are not part of the injection molded base plate of the chip 400. Instead, they are placed on top of the open U-shaped or rectangular channel 402 of the base plate of the chip 400  
10 in a separate lamination step.

Extending from the measurement channel 402 is the waste reservoir 406. The waste reservoir 406 stores the sample solution after its evaluation by the electrochemical sensing in channel 402. The size of the  
15 waste reservoir 406 defines the amount of time one channel can theoretically be used. After the reservoir 406 is filled, the controller can activate the next channel to continue with the detection of glucose in the body fluid of a patient.

20 It is important that the reservoir 406 is large enough to support more than a two-hour measurement. Thus, a chip 400 with twelve channels 402 can run for 24 to 28 hours allowing the patient a convenient change to a new chip (e.g. during his daily morning routines).  
25 However, as will be appreciated by those skilled in the art, reservoirs 406 larger than 5  $\mu$ L (which is enough to support the flow for more than two hours with a flow rate of 30 nL/min) could allow longer runtimes. Thus, the entire chip 400 could last for more than one day. The  
30 number of channels in a device is not limited and may vary depending upon the measurement technique, the length of monitoring period and the size of the device and could exceed 100.

Figures 20a, 20b show two different representative  
35 embodiments of the waste reservoir 406, 406a. Waste reservoir 406 is a meander-shaped extension of the

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channel 402. While easy to manufacture, such a design can build up high back pressure in the system, due to the weaker capillary force, then the design of waste reservoir 406a and is not as space efficient.

5       The column filled reservoir 406a has advantages compared to the meander-shaped reservoir 406 regarding the size and the integration into a 12-channel chip 400a.

It could be deeper than the reservoir 406 while producing a higher capillary force to take up the  
10       extracted and evaluated sample fluid from channel 402a. In principle, waste channel 406a resembles a plurality of capillaries with a cross-section for example of  $400\text{ }\mu\text{m} \times 400\text{ }\mu\text{m}$  compared to only one capillary with the same cross-section in the design of waste reservoir 406. The  
15       shape of the columns is not limited to simple squares. They could be formed as circles, pentagons, hexagons, octagons or the like. Concerning the flow characteristics inside a column field the hexagon shaped columns shown are the most preferred version. The size and geometry of  
20       the channels and reservoirs may vary.

The base plate of the chip 400 can be produced in a molding process, which is optimized for the small features and structures. A wide variation of plastics can be used for the base plate of the chip 400 (e.g.  
25       polystyrene, polycarbonate, polymethymethacrylate, polyester, and others). Preferred polymers are polycarbonates. These allow a subsequent laser finishing to generate a secondary micro- or nano-structure (e.g., any desired patterns or other finishing can be formed in  
30       the micro-channel). Polystyrene shows preferable characteristics in the lamination process. Thus polycarbonate could be used as the lower laminate and polystyrene used as the upper.

The chip 400 itself consists of three major part a)  
35       the chip base plate with all the fluidic elements described above, b) the electrochemical detection system

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screen printed on a polymer foil, and c) the sampling hub with extraction needle. These elements are described in the foregoing sections.

5 Normal lamination processes use foils coated with a pressure sensitive or hot melt adhesive to join a foil to a substrate or another foil. Such a standard process may present problems in conjunction with the described device. First, foil is needed which is suitable for the printing process. This is difficult with any pressure  
10 sensitive adhesive due to problems within the printing equipment. Such problems can be addressed with a foil coated with a hot melt adhesive, where the adhesive becomes tacky only at an elevated temperature (e.g. 80 °C). The deposition of ink to print the electrodes and  
15 other structures is quite easy with this system but it can present problems during the lamination step. The glue layer becomes substantially liquid at elevated temperatures with the consequence that the printed structure loses shape and gets stretched and deformed.  
20 Such deformation is not only a cosmetic problem for an electrode, it changes the electrode surface (which is directly proportional to the response signal) as well as the internal resistance of the material and the electro-catalytic properties.

25 Apart from the foregoing problems, there is the additional problem of glue entering and clogging or misshaping the channel. For the chip described above, the most advantageous process is an adhesive-free thermal bonding process of the pre-printed foil to the chip base  
30 plate. The bonding happens at an elevated temperature with a stamping tool or a hot roller press. The temperature is close to the glass transition temperature ( $T_g$ ) of the polymer thus the low molecular weight portion of the polymer will become mobile and tacky while the  
35 high molecular weight portion of the polymer still supports the integrity of the foil or film. The low

- 45 -

molecular weight portion of the polymer will bond both pieces (base plate and foil with electrodes) together, additionally it will follow the shape of the printed electrodes, which can be between 5 and 30  $\mu\text{m}$  thick.

- 5 Therefore, one does not see a leakage between base plates and printed areas. Ideal bonding is achieved with the same thermoplastic polymers such as polystyrene on polystyrene or polycarbonate on polycarbonate. However, with the right regime and temperature/pressure
- 10 combination polycarbonate can be bonded on polystyrene as well. But duro-plastic (non thermoplastic) materials are not suitable for such a process.

#### 4. Channel Switching Mechanism

- 15 The switching mechanism 404 allows the replacement of a used electrochemical system after a predetermined period of time. Generally, an electrochemical system is prone to fouling due to precipitation or adsorption of proteins or other species on the electrode surface.

- 20 In the past, many different systems have been developed to overcome this problem. A classical example for such a system is the mercury-drop-electrode, where the mercury provides the electrode surface but every second the drop of mercury is replaced. Thus, the system
- 25 never undergoes a continuous fouling process of a solid-state electrode. The renewal of the electrode surface is the most efficient but also the most dramatic strategy to avoid fouling. However, this same concept is used with all current disposable diagnostic glucose strips, where
- 30 the test strip is disposed after a relatively short working time (5 to 15 seconds depending on the device).

- A different strategy is the protection of the electrode by a semi permeable membrane, which allows the analyte to pass through to the electrode but not a large
- 35 protein (or a red blood cell). This concept is adopted in classical oxygen electrodes.

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The chip 400 uses a mixture of both concepts: a plurality of electrodes in different channels as well as an electrode protected by screen printable membrane (see German patent DE10052066.9 and related International  
5 patent application PCT/EP01/12073). This allows the sensor electrodes to operate for several hours before the fouling shows a significant effect on the results and before the controller is switching to the next channel.

Figure 21 shows a schematic diagram of the  
10 switching mechanism 404. At the beginning of a measurement, all channels 402 are filled with air and the entrance of fluid into the channel 402 from the distribution depression 416 is blocked due to the gas back pressure. The switching system 404 includes  
15 electrodes 422 separate from the electrochemical detector electrodes 418, 420. The switching electrodes 404 allow local heating of a very small area on the thin foil such as polystyrene 424 overlying a well 426. The well 426 is in fluid flow communication with the end of  
20 the reservoir 406 except that the fluid flow is normally blocked by the foil 424. Due to this blockage, air cannot escape from the channel 402 and fluid from the distribution depression 416 cannot flow into the channel 402. Upon heating of the foil, this blockage is removed  
25 thus allowing the fluid to enter the channel.

The temperature at the center point of the overlying electrodes 422 (i.e., overlying the foil 424 above the well 426) very quickly rises over the glass transition  
temperature ( $T_g$ ) of the foil in response to an application  
30 of a small current in the range of some 1/1000 Ampere with the effect that the foil 424 forms an opening above the hydrophobic well 426 allowing air in the channel 402 and reservoir 406 to vent into the well 426. In this event, the gas pressure in the channel 402 is relieved  
35 and fluid can flow from the distribution depression 416



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and into the channel 402.

The hydrophobicity of the well 426 prevents the sample fluid from flowing into the well 426 and leaking out of an opening formed in the film by reason of the energized electrodes 404. After the waste reservoir 406 is filled completely, the controller opens a subsequent channel 402 by energizing its switching electrodes 422 to permit air to escape from the channel 402 and for the subsequent channel 402 to fill with body fluid. It should be appreciated that other methods for switching between channels and controlling fluid flow could be used and the invention is not limited by the above method

15

#### 5. Flow Rate Detection

To assure a safe operation, it is preferable the controller determine if the waste reservoir 406 is filled, is still filling at a constant rate, or if the needle 410 has dislodged from the cutaneous layer of the patient and the continuous sample stream has stopped. To achieve this, the waste reservoir 406 can be equipped with two electrodes (not shown) at the top and the bottom of the reservoir. Not in electrical contact with the sample fluid, these electrodes could measure the change in capacitance while the air inside the structure is slowly exchanged with sample fluid. The rate of the capacitance change will be directly proportional to the flow rate of the sample fluid and allow a close monitoring of the ongoing extraction.

#### 6. Semi-Continuous Monitoring Alternative

An alteration of the foregoing is the scheduled measurement of discrete glucose values. With this embodiment the waste reservoir is just large enough to replace the dead volume of the needle and sampling module

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but instead of for example twelve channels the chip contains a higher number of channels (e.g. 72 or 144 channels). These structures would support a 12-hour or 24-hour measurement cycle with a discrete glucose test very 10 minutes. An advantage of the semi-continuous method would be that it would not be necessary to use a cross-linked hydrogel due to the fact that migration of the reagents into the fluid sample over time would no longer an issue. Consequently, conventional enzyme inks such as disclosed in US5708247 could be used.

It will be appreciated by those skilled in the art that whilst some of the potential embodiments of the inventive concepts disclosed herein have been described in greater detail, there are many different variations and modifications to these possible. For example, devices in accordance with the invention may measure the concentration of analytes other than those in bodily fluids.

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## Claims:

1. A device for measuring the concentration of an  
5 analyte in a fluid, comprising a support member and a  
plurality of analyte sensing means provided thereon for  
measuring said concentration, wherein said device further  
comprises a plurality of conduits such that each of said  
sensing means has a conduit associated therewith, said  
10 conduits serving in use to direct said fluid to  
respective sensing means.
2. A device as claimed in claim 1 comprising a  
penetration member, preferably integrally formed,  
15 sufficiently short only to penetrate the cutaneous layer  
of skin, most.
3. A device as claimed in claim 1 or 2 comprising at  
least one microchannel for conveying a interstitial fluid  
20 to one or more of the sensing means.
4. A device as claimed in claim 1, 2 or 3 wherein at  
least some of said sensing means are the same or are for  
measuring the same analyte.
- 25 5. A device as claimed in claim 4 which is arranged  
such that in use the plurality of sensing means take  
measurements of the fluid at different times to enable  
monitoring of the concentration of the analyte in the  
~~fluid over a period of time.~~
- 30 6. A device for measuring the concentration of an  
analyte in a fluid, comprising a support member, analyte  
sensing means provided thereon for measuring said  
concentration and a microchannel in fluid communication  
35 with said analyte sensing means for conveying said fluid  
to said sensing means.

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7. A device as claimed in claim 6 wherein the analyte sensing means comprises one or more reagents to react with said analyte and thereby give a measurable output.
- 5
8. A device as claimed in claim 7 wherein the reagent or at least one of the reagents is provided on a wall of the microchannel.
- 10
9. A device for measuring the concentration of an analyte in a fluid, comprising a support member and a microchannel provided on said support member for conveying said fluid across the support member, wherein one or more reagents for reacting with said analyte is
- 15
- coated on a wall of the microchannel.
10. A device as claimed in claim 9 comprising means for measuring the output of a reaction between the analyte and the reagent.
- 20
11. A device as claimed in any preceding claim comprising integral fluid extraction means for extracting a body fluid from a human or animal body fluid.
- 25
12. A device for measuring the concentration of an analyte in a fluid, comprising a support member, means provided on the support member for sensing said analyte, wherein said support member further comprises fluid
- 30
- extraction means arranged to direct fluid to said analyte sensing means via a conduit on the support member.
13. A device as claimed in claim 12 wherein said conduit comprises a microchannel.
- 35
14. A device as claimed in claim 11, 12 or 13 wherein said fluid extraction means comprises a skin penetration

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member.

15. A device as claimed in claim 14 wherein said skin  
penetration member is of such a length that it penetrates  
5 only the cutaneous layer of skin, not the sub-cutaneous  
layer.

16. A device as claimed in claim 15 wherein said skin  
penetration member is adapted to extract interstitial  
10 fluid.

17. A device for measuring the concentration of an  
analyte in a bodily fluid, comprising skin penetration  
means, said penetration means being sufficiently short so  
15 as to penetrate only the cutaneous layer of skin without  
penetrating the sub-cutaneous layer, and a support  
member, said support member comprising an analyte sensing  
site and a microchannel for conducting said body fluid  
from said penetration member to said analyte sensing  
20 site.

18. A device as claimed in claim 17 wherein said  
penetration means is integral with said support layer.

25 19. A device as claimed in any of claims 14 to 18  
comprising a microneedle for extracting interstitial  
fluid.

20. A device as claimed in any preceding claim which is  
30 adapted to measure glucose concentration.

21. A device as claimed in any preceding claim  
comprising analyte sensing means including a mediated  
amperometric enzyme electrode.

35

22. A device as claimed in claim 21 adapted to utilise

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ferrocene-mediated electron transfer from a glucose-oxidase catalysed reaction.

23. A device as claimed in any preceding claim  
5 comprising a plurality of analyte sensing means.

24. A device as claimed in any preceding claim which is arranged to measure an analyte concentration directly.

10 25. A device as claimed in any preceding claim which is suitable for attachment to the skin of a subject for measuring the concentration of an analyte in blood or interstitial fluid of the subject.

15 26. A device as claimed in claim 25 comprising means for attaching the device to the skin.

27. A device for measuring the concentration of a given analyte in a body fluid comprising means for attaching  
20 the device to the skin of a subject and a plurality of sensing means for making a plurality of measurements of said concentration over a period of time.

28. A device as claimed in any preceding claim arranged  
25 to take measurements at predetermined intervals.

29. A device as claimed in claim 28 comprising flow control means for influencing the flow of fluid to the  
sensing means.

30

30. A device as claimed in claim 29 wherein the flow control means comprises a hydrophobic gate disposed in a microchannel.

35 31. A device as claimed in claim 30 comprising means for changing the hydrophobicity of said gate.

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32. A device as claimed in claim 31 comprising a hydrophilic base material covered by a hydrophobic material and further comprising means for exposing said hydrophilic base material.

5

33. A device as claimed in claim 30 comprising pumping means arranged to apply a pressure to liquid in said microchannel sufficient to breach said hydrophobic gate.

10 34. A device as claimed in any preceding claim comprising a plurality of analyte sensing means, said device being arranged to direct fluid sequentially to each of the sensing means.

15 35. A device as claimed in claim 34 comprising or being adapted to interface with, control means for generating signals to control said direction of the fluid to the sensing means.

20 36. A device as claimed in claim 35 comprising said control configured so as to allow a user to specify when a measurement is to be made.

25 37. A device as claimed in claim 35 or 36 comprising colorimetric transduction means.

30 38. A device as claimed in any of claims 34 to 37 comprising a common fluid collection region arranged so as in use to be in fluid communication with a body fluid to be measured.

35 39. A device as claimed in claim 38 wherein each of said sensing means is in selective fluid communication with said common collection region.

40. A device for making a plurality of measurements of

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the concentration of an analyte in a fluid comprising a common sample collection site in fluid communication with the fluid to be measured and a plurality of sensing means for measuring said concentration.

5

41. A device as claimed in any of claims 34 to 40 comprising a sensing means for making a substantially continuous measurement of the concentration of a substance.

10

42. A device for measuring the concentration of an analyte in a fluid comprising a sensing means for measuring said concentration substantially continuously and calibration sensing means for performing at least one calibration measurement.

15

43. A device as claimed in claim 42 comprising a plurality of individual sensing means to allow a series of periodic calibration measurements to be performed.

20

44. A device as claimed in claim 44 wherein the continuous sensing means and calibration sensing means are provided on a common base member.

25 45. A device as claimed in claim 44, wherein said common base member comprises a patch or the like for attachment to the skin of a user.

30 46. A device as claimed in any of claims 42 to 45 comprising flow control means associated with fluid conduits for conducting fluid to the calibration sensing means.

35 47. A device as claimed in claim 46 wherein said flow control means comprises an electro-osmotic pumping arrangement.



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48. A device as claimed in any of claims 42 to 47 which is suitable for attachment to a user's skin.

49. A device as claimed in any of claims 42 to 48  
5 wherein said calibration sensing means comprises a single use sensor.

50. A device as claimed in any of claims 42 to 49  
10 wherein said calibration sensing means is downstream of the continuous sensing means.

51. A device as claimed in any of claims 42 to 50 comprising one or more microchannels for conveying the fluid to be measured to the sensing means.  
15

52. A device as claimed in any preceding claim which is suitable for attachment to a user's skin comprising transfer means for transferring said a body fluid from the user's body to a sensitive part of the device.  
20

53. A device as claimed in claim 52 wherein said transfer means is arranged to transfer a fluid to the upstream end of a microchannel.

25 54. A device as claimed in claim 52 or 53 wherein said transfer means comprises a needle.

55. A device as claimed in claim 54 wherein said needle  
~~is a microneedle of such a length that it penetrates the~~  
30 cutaneous layer of skin, but not the sub-cutaneous layer.

56. A device as claimed in claim 54 or 55 wherein the needle comprises an aperture on a side surface thereof.

35 57. A device as claimed in any of claims 52 to 56 wherein said transfer means is invasive or semi-invasive,

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the device further comprising means for applying pressure to the user's skin in the region of penetration by said transfer means.

5. 58. A device as claimed in any preceding claim comprising display means for displaying the concentration of an analyte being measured.

59. A device as claimed in claim 58 wherein said  
10 display means is separate from the rest of the device and receives data therefrom by telemetry.

60. A device as claimed in any preceding claim comprising means for administering a substance to a user  
15 on the device on the basis of a measured analyte concentration.

61. An device for administering a substance to a user comprising a measuring device for measuring the  
20 concentration of an analyte in a bodily fluid from said user, said measuring device comprising a plurality of analyte sensing means and at least one conduit, preferably a microchannel, for conveying said body fluid to at least one of the sensing means; the apparatus  
25 further comprising administration means for administering said substance to said user on the basis of said measurement of concentration.

62. ~~A device as claimed in claim 61 wherein the~~  
30 measuring device comprises a self-adhesive patch for attachment to the skin of a user.

63. A device as claimed in claim 61 or 62 wherein said administration means is integrated with the measuring  
35 device.

64. An device as claimed in claim 63 wherein the

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administration means comprises a reservoir on or in the measuring device for dispensing the substance.

65. An device as claimed in claim 64 further comprising  
5 a pump for removing said substance from the reservoir.

66. An device as claimed in claim 65 comprising a single pump for both administering said substance to a user's body and for drawing out blood or interstitial fluid, to  
10 a sensing means for making an analyte concentration measurement.

67. A device as claimed in any preceding claim comprising electro-osmotic flow control means comprising  
15 a plurality of driving electrodes, said driving electrodes being provided substantially on one side of a channel.

68. A device as claimed in claim 67 wherein said driving  
20 electrodes extend circumferentially around an arcuate wall of the channel.

69. A device as claimed in any preceding claim comprising a main support member and a second substrate  
25 laminated to the main support member, said second substrate layer having one or more electrodes formed thereon.

70. A device as claimed in claim 69 wherein said second  
30 substrate is arranged to close a channel provided on the main support member.

71. A device for measuring the concentration of an analyte in a fluid comprising a support member having an  
35 open channel on a surface thereof the support member; and a second layer laminated onto said support member so as

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to close said channel.

72. A device as claimed in claim 71 wherein said channel is a microchannel.

5

73. A device as claimed in claim 71 or 72 comprising an integrated skin penetration member is provided which is also open on one side.

10 74. A device as a claimed in claim 73 wherein said skin penetration member is arranged so that upon insertion into skin, the skin itself effectively forms a wall of

75. A device for obtaining fluid through human or animal  
15 skin, comprising a skin penetration member having at least one longitudinal side open, the other sides being arranged so as effectively to cause the penetrated skin to act as the remaining longitudinal side of the member when the penetration member is inserted into the skin.

20

76. Apparatus for measuring an analyte optically comprising a device as claimed in any preceding claim for measuring said analyte and a separate test meter including light sensitive means, wherein said measuring  
25 device and said test meter are arranged such that in use the light sensitive means views said analyte along a channel of the measuring device.

77. Apparatus for measuring the light from an assay  
30 comprising an elongate conduit portion along which a sample fluid is drawn in use and a light sensitive means arranged to be sensitive to light coming substantially from the longitudinal axis of said conduit portion.  
78. A disposable skin patch comprising an elongate  
35 conduit portion along which a sample fluid is drawn in use and a moulded plastics lens over said conduit

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portion.

79. A test meter for use with disposable skin patch as claimed in claim 78 comprising a light sensitive element arranged, in use, to sit over said lens when the meter is placed over the patch.

80. A method of measuring the amount of an analyte in a sample of liquid comprising providing an electrochemical measuring device having a sensor electrode within a microchannel, introducing said sample liquid into said microchannel, and measuring an aggregate current passed by said electrode to give an indication of said amount of analyte in the sample.

81. An electrochemical device for measuring the amount of an analyte in a sample of liquid comprising a sensor electrode disposed within a microchannel and means for measuring an aggregate current passed by said electrode in use to give an indication of said amount of analyte in the sample.

82. A method of making a device for measuring the concentration of a substance in a liquid comprising forming a first channel in a substrate material, fitting said channel at least partially with an electrically conductive material, and forming a second channel so as to intersect said first channel, thereby forming two conductive portions on respective opposite sides of the second channel.

83. A method of fabricating a device for measuring the concentration of an analyte in a fluid comprising providing a support member, forming an open channel on a surface of the support member and laminating a second layer onto said support member so as to close said

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channel.

84. An apparatus for obtaining and analyzing a sample of a body fluid from a skin layer, said apparatus  
5 comprising:

a support member;

a channel formed on said support member;

said channel having an entrance for receiving said body fluid;

10 an analyte sensor carried on said support member and positioned to sense an analyte in said fluid within said channel;

a flow controller for influencing a flow of fluid within said channel to said sensor.

15

85. An apparatus according to claim 84 further comprising a fluid extraction member for directing fluid from said skin layer to said entrance.

20 86. An apparatus according to claim 85 further comprising a flow restrictor between said fluid extraction member and said channel.

87. An apparatus according to claim 84 further  
25 comprising a waste reservoir in fluid flow communication with said channel.

88. An apparatus according to claim 84 wherein said flow controller includes a flow inhibitor in said channel  
30 inhibiting flow of said body fluid into said channel and a reliever for permitting discharge of said inhibitor from said channel to permit said fluid to flow into said channel.

35 89. An apparatus according to claim 88 wherein said flow inhibitor is an inhibitor fluid in said channel and said

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reliever includes a switch for permitting selective release of said inhibitor fluid from said channel.

90. An apparatus according to claim 89 wherein said  
5 switch includes a membrane blocking passage of said inhibitor fluid from said channel and selectively controllable electrodes for removing said membrane.

91. An apparatus according to claim 90 wherein said  
10 membrane is selected to be at least partially destroyed by energization of said electrodes.

92. An apparatus according to claim 89 wherein said  
15 inhibitor fluid is air and said switch vents said air from said channel.

93. An apparatus according to claim 84 wherein said flow controller includes a hydrophobic gate.

20 94. An apparatus according to claim 93 wherein said hydrophobic gate includes a hydrophobic length of said channel.

95. An apparatus according to claim 84 wherein said flow  
25 controller includes a bubble disposed in said channel with a vent to said channel to permit selective discharge of said bubble from said channel.

96. An apparatus according to claim 84 wherein said flow  
30 controller further comprises a pumping mechanism for urging fluid flow past said flow controller.

97. An apparatus according to claim 96 wherein said  
35 pumping mechanism is an electro-osmotic pump.

98. An apparatus according to claim 96 wherein said

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pumping mechanism is automatically controllable by a controller.

99. An apparatus according to claim 96 wherein said  
5 pumping mechanism is selectively controllable by a user.

100. An apparatus according to claim 84 wherein:  
said channel is one of a plurality of channels each  
formed on said support chamber by forming channel  
10 defining walls on a said support member and each having  
an entrance for receiving said body fluid;  
said analyte sensor is one of a plurality of analyte  
sensors each associated with a respective one of said  
plurality of channels.

15  
101. An apparatus according to claim 100 wherein said  
controller operates to direct flow into a subsequent one  
of said plurality of channels after a residence time of  
said body fluid in a preceding one of said plurality of  
20 channels.

102. An apparatus according to claim 100 wherein each of  
said entrances is connected to a common source of said  
fluid.

25  
103. An apparatus according to claim 100 wherein said  
plurality of channels and associated sensors include:  
at least a first continuous sensor for sensing a  
concentration of an analyte in fluid within said channel  
30 associated with said continuous sensor; and  
at least a first calibration sensor for performing  
as least one calibration measurement of a concentration  
of an analyte in fluid within said channel associated  
with said calibration sensor.

35  
104. A method of continuously sampling and measuring an



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analyte comprising:

extracting a fluid into a common collection port;  
directing said fluid along a channel towards an  
analyte sensing means;

5 performing at least an analyte measurement on the  
fluid;

switching the flow of fluid from one channel to  
sequentially at least another channel after a certain  
period of time; and

10 performing at least a further analyte measurement.

105. An apparatus according to claim 84 further  
comprising an administrator for administering a substance  
to a user based upon a measuring based upon a sensing of  
15 said analyte sensor.

106. An apparatus for obtaining and analyzing a sample of  
a body fluid from a skin layer, said apparatus  
comprising:

20 a support member;

a channel formed on said support chamber by forming  
channel defining wall on a said support member;

said channel having an entrance for receiving said  
body fluid;

25 said channel having a minute size with a volume less  
than 200 nanoliters;

an analyte sensor carried on said support member and  
positioned to sense an analyte in said fluid within said  
channel.

30

107. An apparatus according to claim 106 wherein said  
channel has a longitudinal dimension less than 10 mm.

108. An apparatus according to claim 106 wherein said  
35 channel has a lateral dimension less than 200  
micrometres.

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109. An apparatus according to claim 108 wherein said lateral dimension is greater than 10 micrometres.
- 5 110. An apparatus according to claim 106 wherein said analyte sensor includes a reagent deposited on said wall for reacting with an analyte of said fluid when said fluid is within said channel.
- 10 111. An apparatus according to claim 106 further comprising a fluid extraction member for directing fluid from said skin layer to said entrance.
112. An apparatus according to claim 111 wherein said  
15 fluid extraction member is a penetration member for penetrating into only a cutaneous layer of said skin layer and said fluid extraction member has a fluid pathway for fluid to flow from said cutaneous layer to said entrance of said channel.
- 20 113. An apparatus according to claim 112 comprising a plurality of said fluid extraction members.
114. An apparatus according to claim 112 wherein said  
25 penetration member is a hollow needle with a bore of said needle defining said fluid pathway.
115. An apparatus according to claim 112 wherein said  
penetration member is a lance having opposing surfaces  
30 defining said fluid pathway and with said fluid pathway open to an exterior of said lance.
116. An apparatus according to claim 111 wherein said  
fluid extraction member includes means for iontophoretic  
35 transfer of fluid from said skin layer to said entrance.

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117. An apparatus according to claim 111 wherein said fluid extraction member includes means for ultra-sonic transfer of fluid from said skin layer to said entrance.
- 5 118. An apparatus according to claim 111 wherein said fluid extraction member includes means for suction-assisted transfer of fluid from said skin layer to said entrance.
- 10 119. An apparatus according to claim 111 further comprising an electric field generator for stimulating the skin layer and urging fluid to flow to said entrance.
- 15 120. An apparatus according to claim 119 wherein said fluid extraction member includes an electrode which comprises a portion of said electric field generator.
- 20 121. An apparatus according to claim 86 further comprising a flow controller for influencing a flow of fluid within said channel to said sensor.
122. An apparatus according to claim 86 wherein:  
said channel is one of a plurality of channels each formed on said support chamber by forming channel  
25 defining walls on a said support member and each having an entrance for receiving said body fluid;  
said analyte sensor is one of a plurality of analyte sensors each associated with a respective one of said  
plurality of channels.
- 30 123. An apparatus according to claim 101 wherein each of said entrances is connected to a common source of said fluid.
- 35 124. An apparatus according to claim 101 wherein said plurality of channels and associated sensors include:

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at least a first continuous sensor for sensing a concentration of an analyte in fluid within said channel associated with said continuous sensor; and

5 at least a first calibration sensor for performing as least one calibration measurement of a concentration of an analyte in fluid within said channel associated with said calibration sensor.

125. An apparatus according to claim 106 wherein said  
10 analyte sensor includes an electro-chemical sensor.

126. An apparatus according to claim 106 wherein said analyte sensor includes a plurality of electrodes.

15 127. An apparatus according to claim 106 wherein at least one of said electrodes is an electrically conductive material deposited in said channel.

128. An apparatus according to claim 106 wherein said  
20 channel is formed in an electrically conductive material with said material comprising at least one of said electrodes.

129. An apparatus according to claim 106 wherein said  
25 analyte sensor includes a colorimetric sensor.

130. An apparatus according to claim 106 further comprising an administrator for administering a substance  
to a user based upon a measuring based upon a sensing of  
30 said analyte sensor.

131. An apparatus for obtaining and analyzing a sample of a body fluid from a skin layer, said apparatus comprising:

35 a support member;  
a channel formed on said support chamber by forming

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channel defining wall on a said support member;

said channel having an entrance for receiving said body fluid;

5       said channel having a reagent deposited on said wall  
for reacting with an analyte of said fluid when said fluid is within said channel.

10       132. An apparatus according to claim 131 wherein said channel has a volume less than 200 nanoliters.

133. An apparatus according to claim 132 wherein said channel has a longitudinal dimension less than 10 mm.

15       134. An apparatus according to claim 132 wherein said channel has a lateral dimension less than 500 micrometres.

20       135. An apparatus according to claim 134 wherein said lateral dimension is less than 200 micrometres.

136. An apparatus according to claim 135 wherein said lateral dimension is greater than 10 micrometres

25       137. An apparatus according to claim 131 further comprising a fluid extraction member for directing fluid from said skin layer to said entrance.

30       138. An apparatus according to claim 137 wherein said fluid extraction member is a penetration member for penetrating into only a cutaneous layer of said skin layer and said fluid extraction member has a fluid pathway for fluid to flow from said cutaneous layer to said entrance of said channel.

35       139. An apparatus according to claim 138 wherein said penetration member is a hollow needle with a bore of said

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needle defining said fluid pathway.

140. An apparatus according to claim 138 wherein said penetration member is a lance having opposing surfaces  
5 defining said fluid pathway and with said fluid pathway open to an exterior of said lance.

141. An apparatus according to claim 131 further comprising a flow controller for influencing a flow of  
10 fluid within said channel to a portion of said channel containing said reagent.

142. An apparatus according to claim 131 wherein:  
said channel is one of a plurality of channels each  
15 formed on said support chamber by forming channel defining walls on a said support member and each having an entrance for receiving said body fluid;  
each of said channels provided with a respective reagent on a defining wall of said channels.

20

143. An apparatus according to claim 142 wherein each of said entrances is connected to a common source of said fluid.

25 144. An apparatus according to claim 142 wherein said plurality of channels includes:

at least a first continuous sensor for sensing a concentration of an analyte in fluid within said channel associated with said continuous sensor; and

30 at least a first calibration sensor for performing as least one calibration measurement of a concentration of an analyte in fluid within said channel associated with said calibration sensor.

145. An apparatus for obtaining and analyzing a sample of  
35 a body fluid from a skin layer, said apparatus comprising:

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- a support member;  
a channel formed on said support chamber by forming  
channel defining wall on a said support member;  
said channel having an entrance for receiving said  
5 body fluid;  
a fluid extraction member for directing fluid from  
said skin layer to said entrance;  
said channel having a minute size with a volume less  
than 200 nanoliters;  
10 an analyte sensor carried on said support member and  
positioned to sense an analyte in said fluid within said  
channel.
146. An apparatus according to claim 145 wherein said  
15 channel has a longitudinal dimension less than 10 mm.
147. An apparatus according to claim 145 wherein said  
channel has a lateral dimension is less than 200  
micrometres.  
20
148. An apparatus according to claim 147 wherein said  
lateral dimension is greater than 10 micrometres.
149. An apparatus according to claim 145 wherein said  
25 analyte sensor includes an electrochemical sensor  
positioned to be in contact with said fluid when said  
fluid is within said channel.
- ~~150. An apparatus according to claim 145 wherein said~~  
30 fluid extraction member is a penetration member for  
penetrating into only a cutaneous layer of said skin  
layer and said fluid extraction member has a fluid  
pathway for fluid to flow from said cutaneous layer to  
said entrance of said channel.
- 35
151. An apparatus according to claim 150 wherein said

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penetration member is a hollow needle with a bore of said needle defining said fluid pathway.

152. An apparatus according to claim 150 wherein said  
5 penetration member is a lance having opposing surfaces defining said fluid pathway and with said fluid pathway open to an exterior of said lance.

153. An apparatus according to claim 145 further  
10 comprising a flow controller for influencing a flow of fluid within said channel to said sensor.

154. An apparatus according to claim 145 wherein:  
said channel is one of a plurality of channels each  
15 formed on said support chamber by forming channel defining walls on a said support member and each having an entrance for receiving said body fluid;  
said analyte sensor is one of a plurality of analyte  
sensors each associated with a respective one of said  
20 plurality of channels.

155. An apparatus according to claim 154 wherein each of  
said entrances is connected to a common source of said  
fluid.  
25

156. An apparatus according to claim 154 wherein said  
plurality of channels and associated sensors include:  
at least a first continuous sensor for sensing a  
concentration of an analyte in fluid within said channel  
30 associated with said continuous sensor; and  
at least a first calibration sensor for performing  
as least one calibration measurement of a concentration  
of an analyte in fluid within said channel associated  
with said calibration sensor.

35  
157. An apparatus according to claim 84 further



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comprising an administrator for administering a substance to a user based upon a measuring based upon a sensing of said analyte sensor.

- 5 158. An apparatus according to claim 84 wherein said analyte sensor includes a light sensor disposed to be sensitive to light coming substantially from a longitudinal axis of said channel.
- 10 159. An apparatus for obtaining and analyzing a sample of a body fluid from a skin layer, said apparatus comprising:
- a support member;
  - a plurality of channels each formed on said support
  - 15 chamber by forming channel defining walls on a said support member;
  - each of said channels having an entrance for receiving said body fluid;
  - a plurality of analyte sensors each associated with
  - 20 a respective one of said plurality of channels, each of said analyte sensors carried on said support member and positioned to sense an analyte in said fluid within said respective one of said plurality of channels.
- 25 160. An apparatus according to claim 159 wherein said channel has a minute size with a lateral dimension less than 500 micrometres.
- 30 161. An apparatus according to claim 159 wherein said channel has a volume less than 200 nanoliters.
162. An apparatus according to claim 161 wherein said channel has a longitudinal dimension less than 10 mm.
- 35 163. An apparatus according to claim 160 wherein said lateral dimension is less than 200 micrometres.

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164. An apparatus according to claim 163 wherein said lateral dimension is greater than 10 micrometres
- 5 165. An apparatus according to claim 159 wherein said analyte sensor includes an electrochemical sensor positioned to be in contact with said fluid when said fluid is within said channel.
- 10 166. An apparatus according to claim 159 further comprising a fluid extraction member for directing fluid from said skin layer to said entrance.
- 15 167. An apparatus according to claim 166 wherein said fluid extraction member is a penetration member for penetrating into only a cutaneous layer of said skin layer and said fluid extraction member has a fluid pathway for fluid to flow from said cutaneous layer to said entrance of said channel.
- 20 168. An apparatus according to claim 167 wherein said penetration member is a hollow needle with a bore of said needle defining said fluid pathway.
- 25 169. An apparatus according to claim 167 wherein said penetration member is a lance having opposing surfaces defining said fluid pathway and with said fluid pathway open to an exterior of said lance.
- 30 170. An apparatus according to claim 159 wherein each of said entrances is connected to a common source of said fluid.
- 35 171. An apparatus according to claim 159 wherein said plurality of channels and associated sensors include:  
at least a first continuous sensor for sensing a

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concentration of an analyte in fluid within said channel associated with said continuous sensor; and

at least a first calibration sensor for performing as least one calibration measurement of a concentration  
5 of an analyte in fluid within said channel associated with said calibration sensor.

172. An apparatus according to claim 159 further comprising an administrator for administering a substance  
10 to a user based upon a measuring based upon a sensing of said analyte sensors.

173. A method of making a device for measuring the concentration of a substance in a liquid, said method  
15 comprising:

forming a first channel in a substrate material;  
fitting said channel at least partially with an electrically conductive material;  
forming a second channel so as to intersect with  
20 said first channel.

174. A method of fabricating a device for measuring the concentration of an analyte in a fluid, said method comprising:  
25 forming a support member;  
forming an open channel on a surface of the support member;  
laminating a layer onto said support member so as to close said channel.

30

175. An apparatus according to claim 174 wherein said laminate layer contains electrodes disposed to be in contact with said open channel.

176. An apparatus according to claim 175 wherein said  
35 electrodes are formed through a screen-printing process.

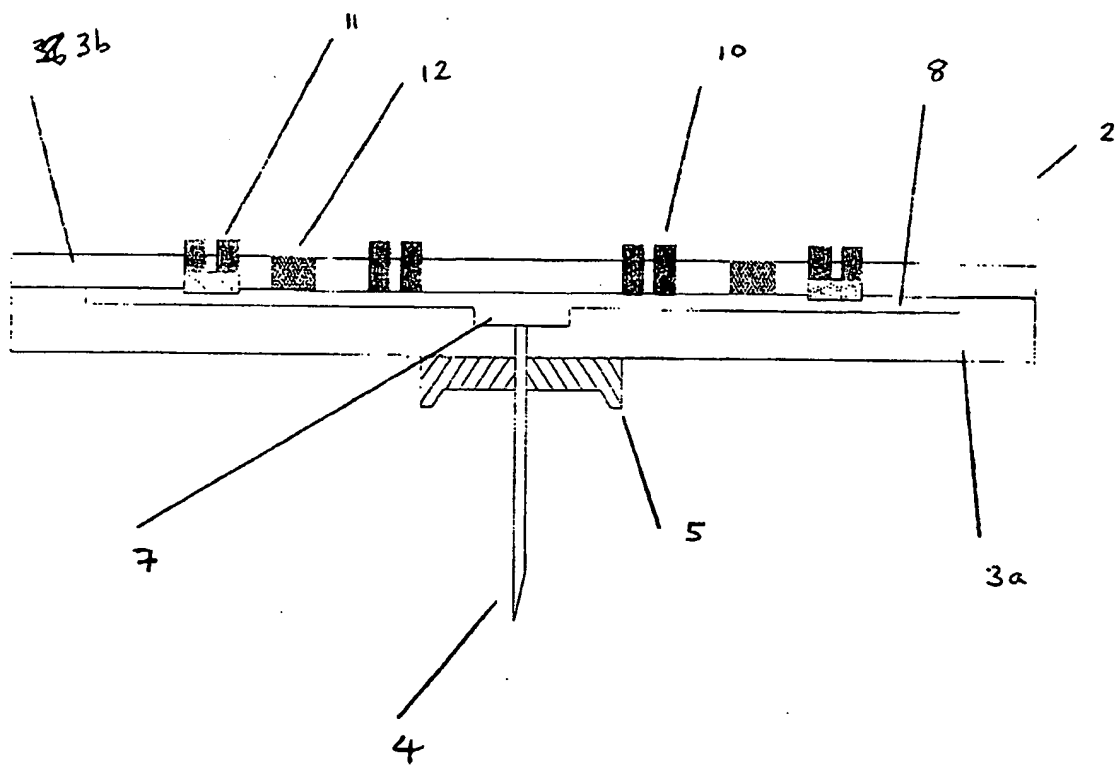


Fig.: 1



Fig 2a

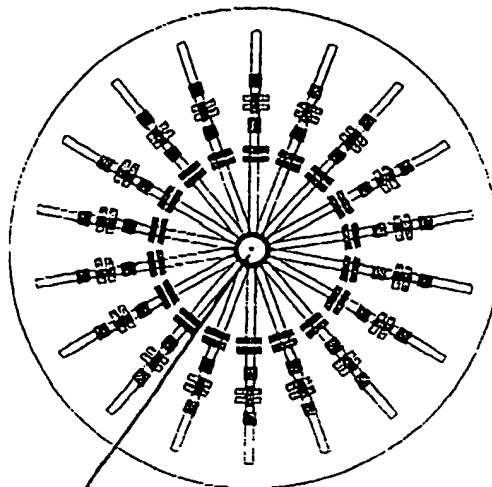


Fig 2b

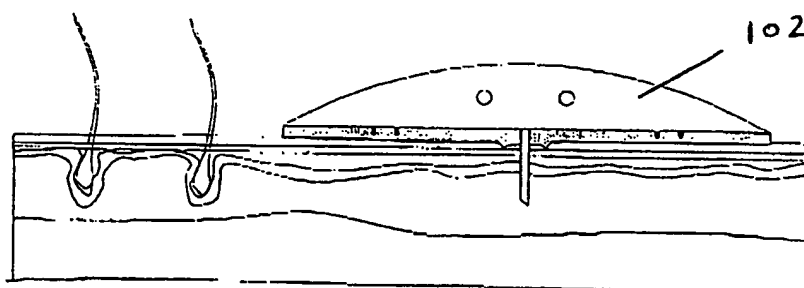


Fig 3

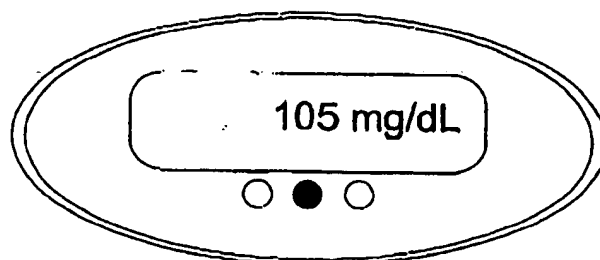


Fig 4

Fig.: 2 to 4

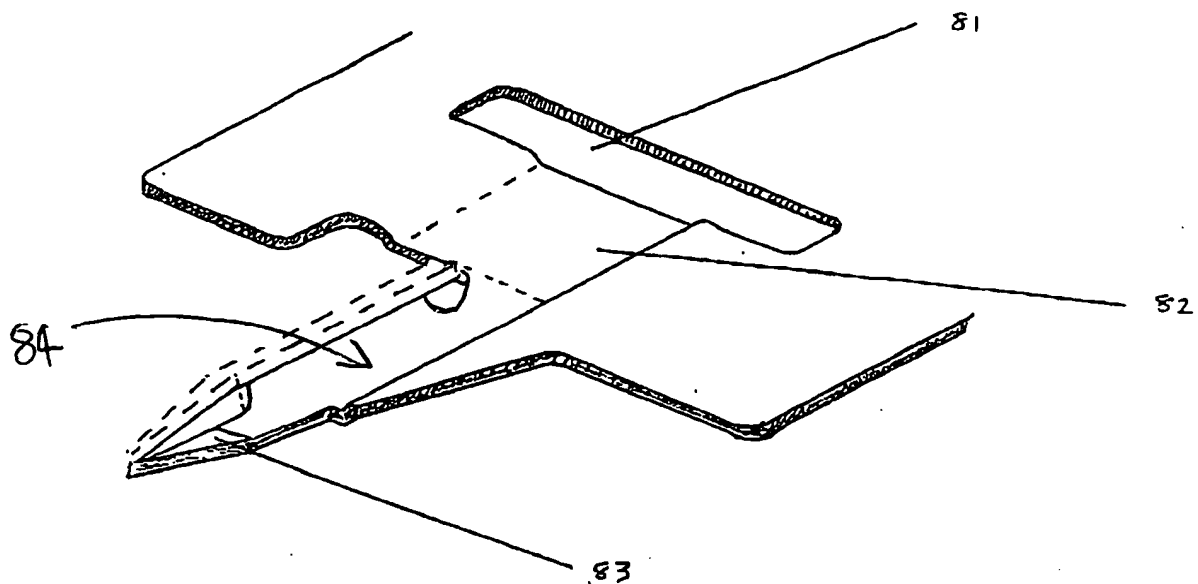


Fig 8i

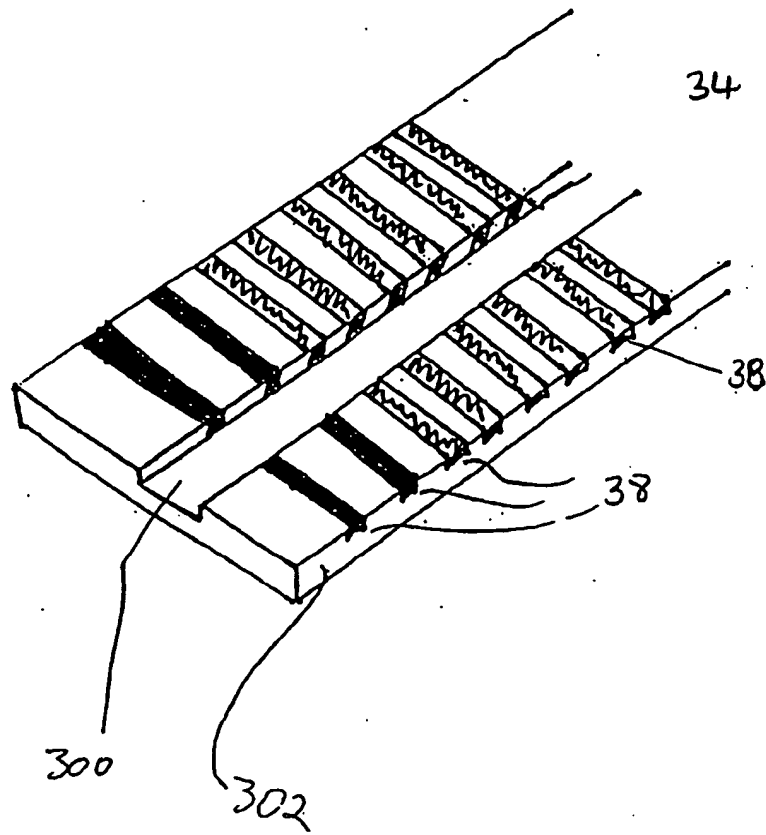


Fig.: 12 a

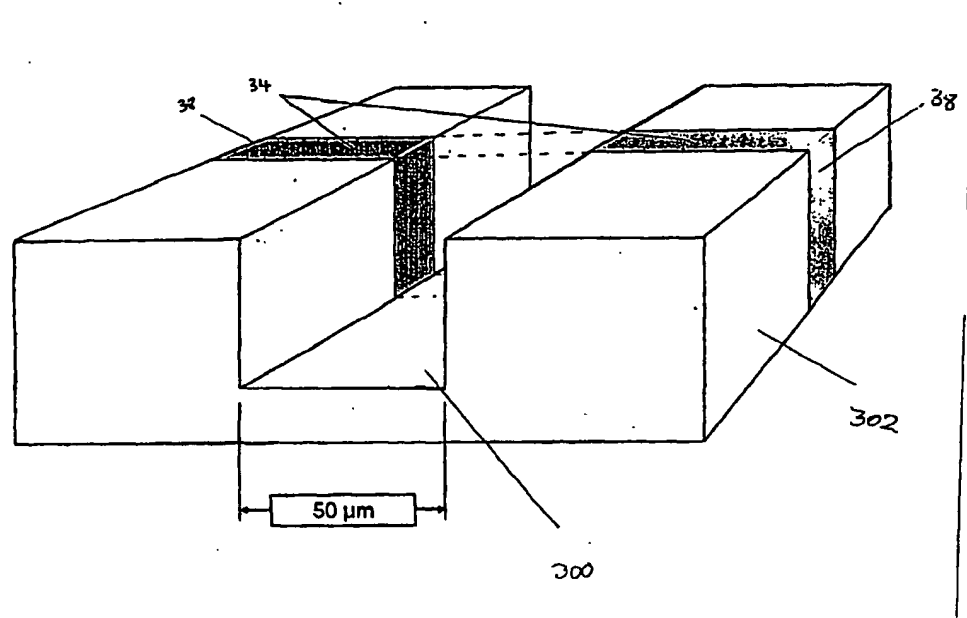


Fig.: 12 b



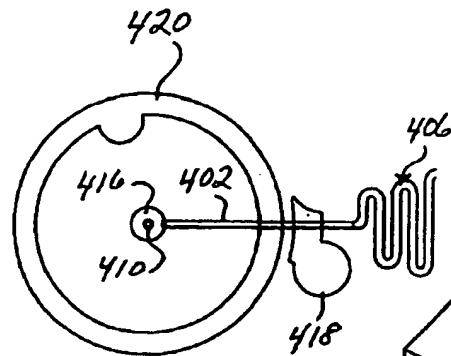


Figure 19

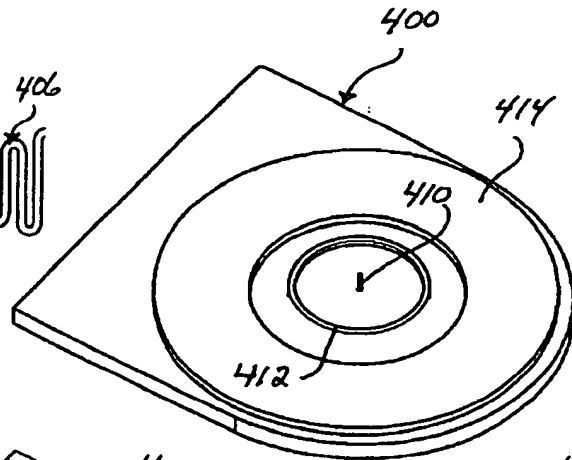


Figure 18

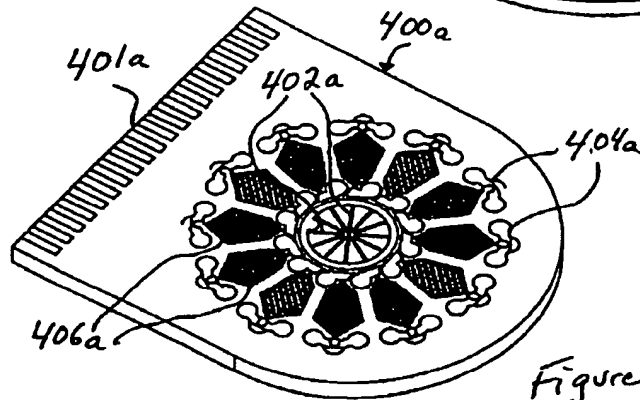


Figure 17b

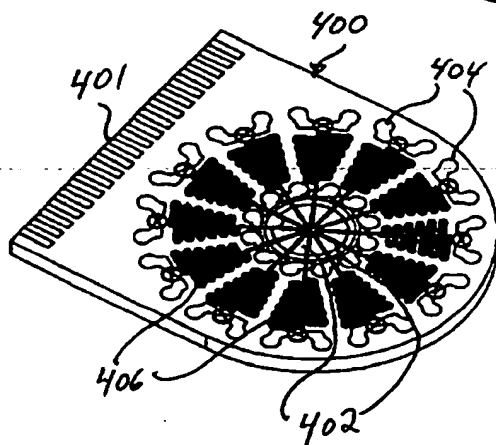
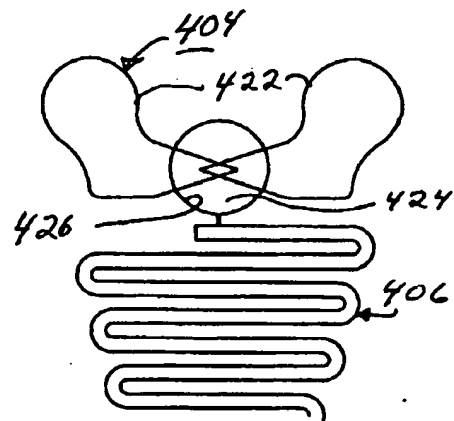
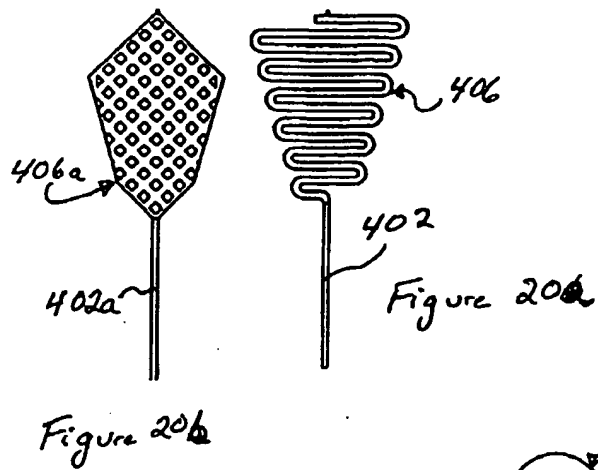


Figure 17a



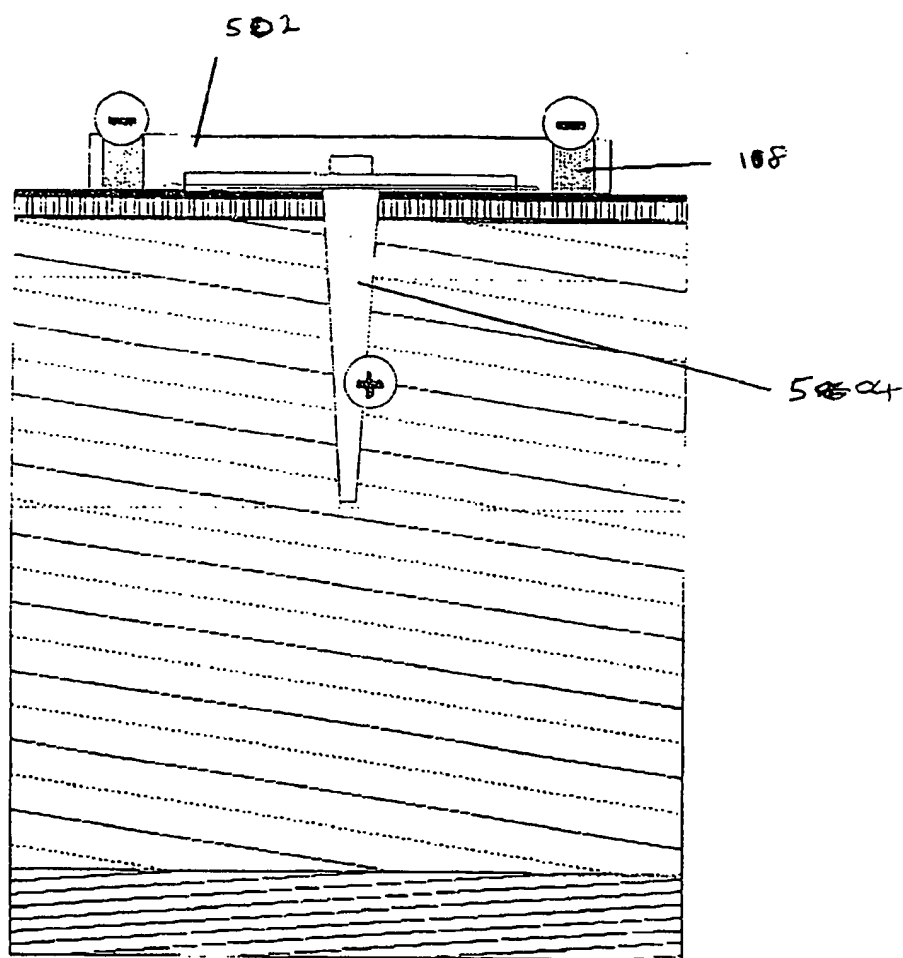
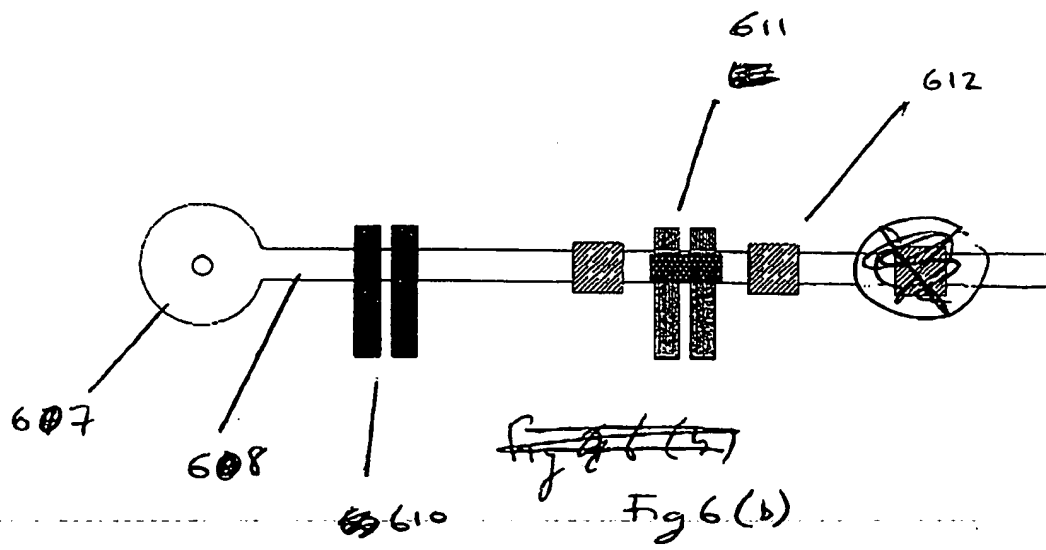
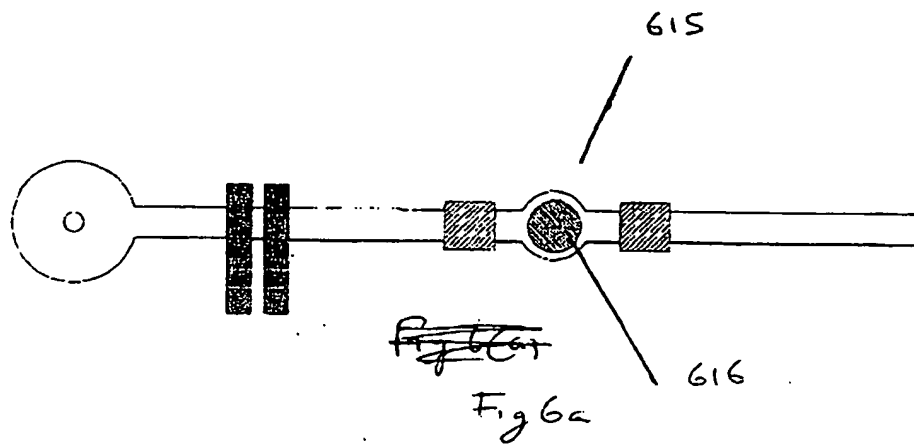
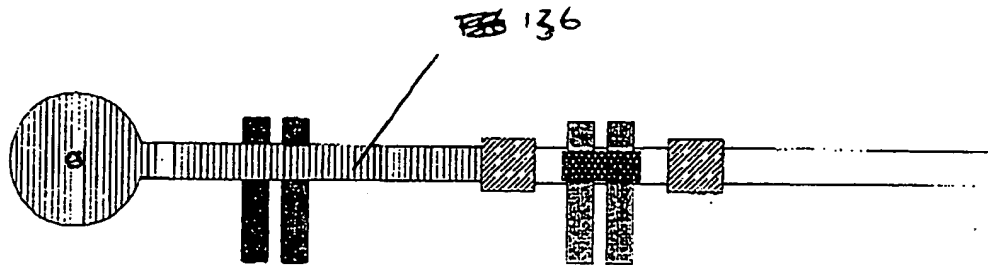
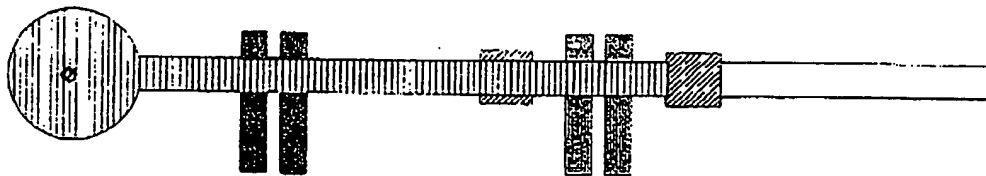


Fig.: 6 S

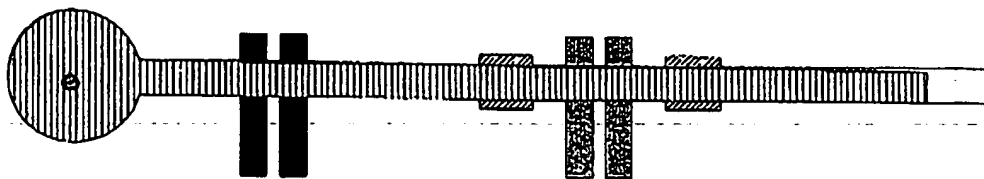




~~Fig 6a~~ Fig 6c

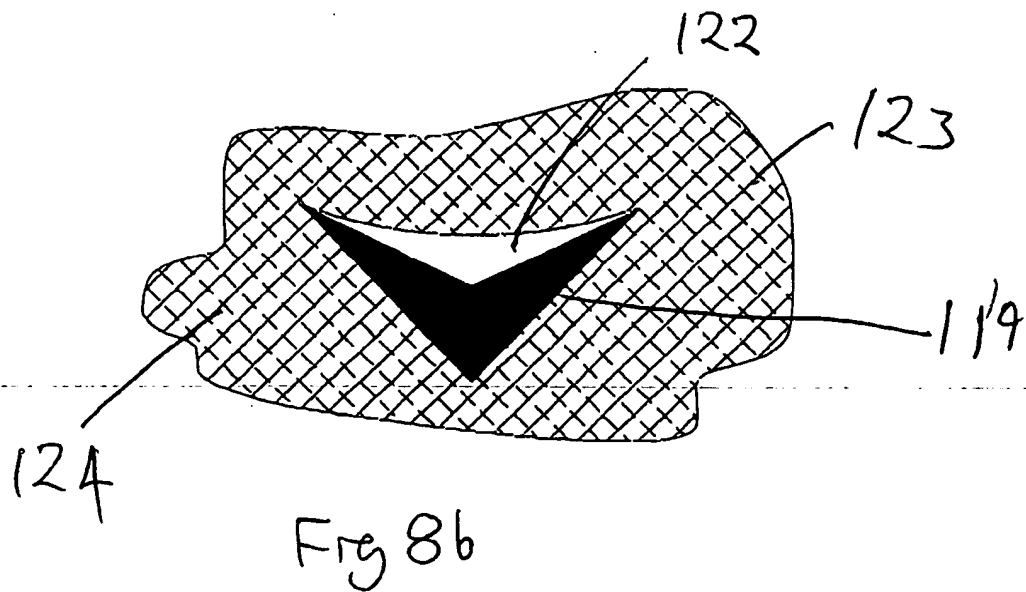
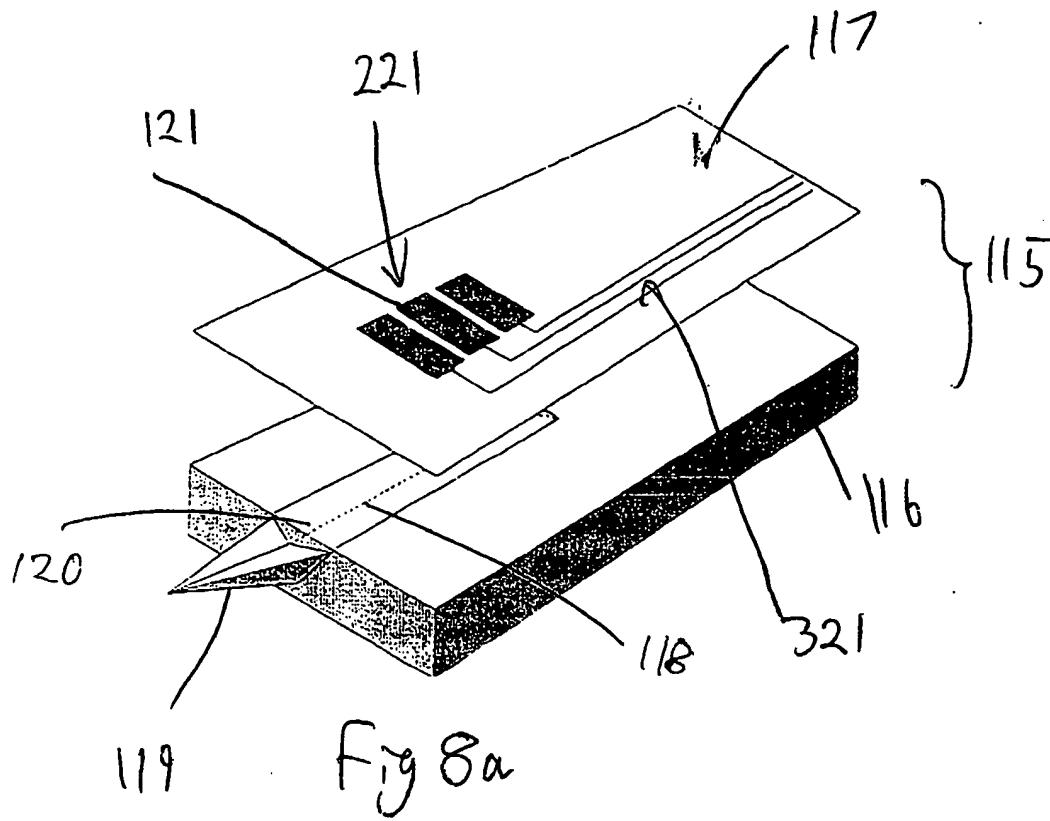


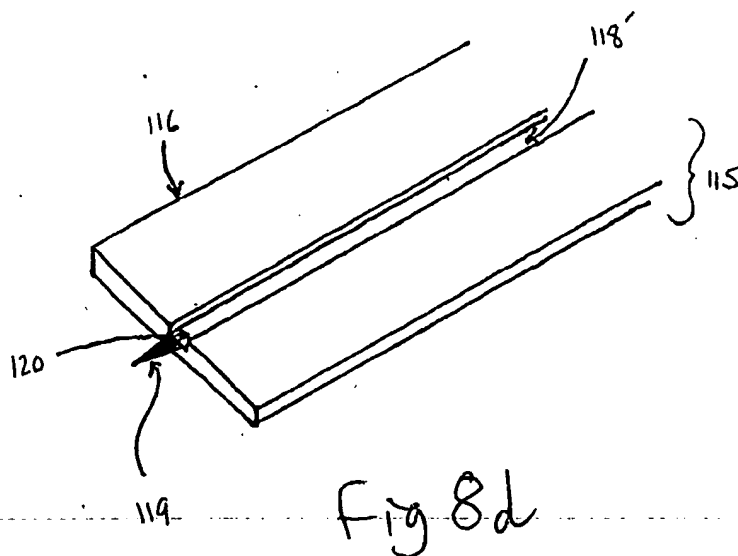
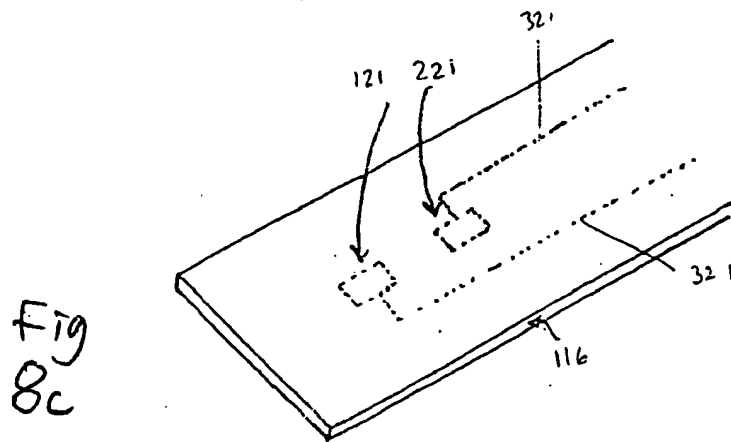
~~Fig 6b~~ Fig 6d



~~Fig 6c~~ Fig 6e

~~Fig 11a to c~~





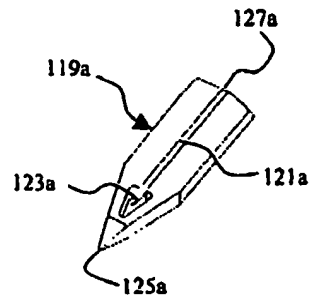


Figure 8e

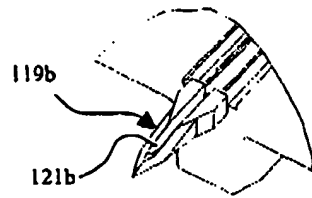


Figure 8f

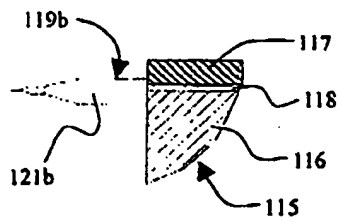


Figure 8g

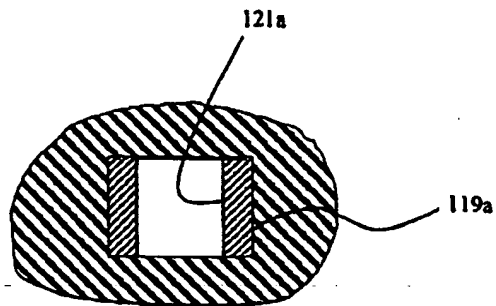


Figure 8h



## INTERNATIONAL SEARCH REPORT

International Application No.

PCT/GB 01/05634

## A. CLASSIFICATION OF SUBJECT MATTER

IPC 7 A61B5/00 A61B5/15 A61B10/00

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 A61B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the International search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	<p>WO 97 19344 A (WILGUS ERIC S ;WARD W KENNETH (US); LEGACY GOOD SAMARITAN HOSPITAL) 29 May 1997 (1997-05-29)</p> <p>page 9, line 17 -page 11, line 7 page 15, line 21 -page 17, line 11; tables 1-3,7-9</p> <p style="text-align: center;">--- -/--</p>	<p>1-5,27, 40,59, 61,100, 122,159</p>

☒ Further documents are listed in the continuation of box C.☒ Patent family members are listed in annex.

## \* Special categories of cited documents:

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- \*O\* document referring to an oral disclosure, use, exhibition or other means
- \*P\* document published prior to the international filing date but later than the priority date claimed

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Date of the actual completion of the international search

26 February 2002

Date of mailing of the international search report

04/03/2002

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Weihs, J

## INTERNATIONAL SEARCH REPORT

Int'l Application No

PCT/GB 01/05634

## C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	<p>WO 00 22977 A (KNOLL MEINHARD) 27 April 2000 (2000-04-27) cited in the application</p> <p>page 3, line 11 -page 9, line 15 page 10, line 20 -page 27, line 24; tables 1-11</p> <p>---</p>	<p>6,12-26, 42-58, 71-74, 80-86, 106-116, 131-139, 145,150, 151, 173-175</p>
A	<p>WO 95 10221 A (KORF JAKOB ;TERWEE THOMAS HENRICUS MARIE (NL)) 20 April 1995 (1995-04-20)</p> <p>abstract page 7, line 3 -page 14, line 32; tables 1,2</p> <p>---</p>	<p>28-33, 84-87, 93-98, 106-108, 131-135, 145-147</p>
A	<p>US 4 873 993 A (MESEROL PETER M ET AL) 17 October 1989 (1989-10-17) abstract</p> <p>---</p>	<p>76,77</p>
P,A	<p>WO 01 64105 A (MOERMAN PIET ;STEINE MATTHIAS (GB); MCALEER JEROME F (GB); INVERNE) 7 September 2001 (2001-09-07)</p> <p>abstract page 10, line 22 -page 15, line 4; tables 7-10</p> <p>-----</p>	<p>1,17-40, 59,61, 75,100, 122,138, 139,145, 150,151, 156,159</p>

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